NATIONAL ADVISORY COMMITTEE FOR AERONAUTICS

TECHNICAL NOTE 2683

SURVEY OF PORTIONS OF THE CHROMIUM-COBALT-NICKEL-

MOLYBDENUM QUATERNARY SYSTEM AT 1200° C

By Sheldon Paul Rideout and Paul A. Beck

University of Notre Dame

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SUMMARY

A survey was made of portions of the chromium-cobalt-nickel-molybdenum quaternary system at 1200°C by means of microscopic and X-ray diffraction studies. Since the face-centered cubic (alpha) solid solutions form the matrix of almost all practically useful high-temperature alloys, the solid solubility limits of the quaternary alpha phase were determined up to 20 percent molybdenum. The component cobalt-nickel-molybdenum, chromium-cobalt-molybdenum, and chromium-nickel-molybdenum ternary systems were also studied. The survey of these systems was confined to the determination of the boundaries of the face-centered cubic (alpha) solid solutions and of the phases coexisting with alpha at 1200°C.

INTRODUCTION

In the development of technologically useful alloys it is usually of considerable help if the phase relationships and solid solubility limits are known. At the Metallurgy Department of the University of Notre Dame, a project has been in progress for some years to determine the phase relationships in alloy systems involving chromium, cobalt, nickel, iron, and molybdenum, the transition elements of greatest importance in high-temperature alloys.

The determination of phase diagrams for systems of four or more components is an extremely laborious task. The problem must be approached in a systematic manner in order to avoid becoming hopelessly lost. The best method of attack is to begin by establishing the phase relationships in systems of two or three components and then continue by adding one new element at a time. The problem of presenting quantitative phase relationships diagrammatically for systems of three or more components necessitates holding one or more thermodynamic variables constant. For example, a ternary phase diagram may be presented as a series of isothermal sections or as a series of sections in each of which the amount of one component is held constant. For a quaternary system, it is necessary to hold both temperature and the amount of one component constant in order to obtain

two-dimensional diagrams. The temperature 1200° C was chosen as that at which an initial isothermal survey could be most profitably made. This temperature is of immediate interest because it lies within the range of solution treatment for most high-temperature alloys now in use and also because here diffusion rates are fast enough to allow equilibrium conditions to be approached in reasonably short annealing periods. At lower temperatures, such as 800° C, the determination of these phase diagrams within extensive composition ranges would be too time-consuming and therefore expensive. Work of this kind is planned only for limited important composition ranges.

Thus far, two reports have been issued covering work done on this project. In the first report (reference 1) the 1200° C isothermal section of the chromium-cobalt-nickel ternary system was presented. The second report (reference 2) gave a survey of the chromium-cobalt-nickeliron quaternary system at 1200° C in the composition ranges near the face-centered cubic solid-solution phase. This report also presented a study of some features of the chromium-cobalt-nickel and the chromium-cobalt-iron ternary systems at lower temperatures.

The present report presents a survey of portions of the chromium-cobalt-nickel-molybdenum quaternary system at 1200° $\overline{\text{C}}$. The face-centered cubic (alpha) solid solutions have, by far, the greatest practical importance in high-temperature alloys. For this reason, the solid solubility limits of the quaternary alpha phase were determined up to 20 percent molybdenum.

It was also necessary to investigate the component cobalt-nickel-molybdenum, chromium-cobalt-molybdenum, and chromium-nickel-molybdenum ternary systems, since these systems had not been adequately explored by previous investigators. (See "Literature Survey" for discussion of previous studies of the binary and ternary systems comprising the chromium-cobalt-nickel-molybdenum quaternary system.)

At the beginning of this work it became apparent that a definite and consistent nomenclature for phase designation would have to be adopted in order to avoid confusion and contradiction. For example, the iron-chromium sigma phase is isomorphous with the chromium-cobalt gamma phase and the iron-molybdenum zeta phase. In order to avoid contradiction in ternary and quaternary systems of these elements, where the isomorphous phases form uninterrupted solid solutions, it was decided to give all these phases the same designation. The designation sigma phase was selected as one most-generally associated with this particular structure. Since this work began with a nonferrous system, the face-centered cubic solid solutions in the systems involving chromium, cobalt, nickel, and molybdenum have all been designated as alpha phase. For the sake of consistency, this nomenclature was extended even to the chromium-cobalt-nickel-iron quaternary system, although this phase in ferrous

systems has been customarily referred to as gamma. On the other hand, the body-centered cubic solid solutions, which are usually named alpha in ferrous systems, are here referred to as epsilon phase in conformity with the nomenclature used for the chromium-rich alloys in the chromium-cobalt binary system. In the cobalt-molybdenum binary system, the intermediate phase which coexists with alpha at 1200° C has been earlier designated as epsilon. In the present investigation, this phase was renamed mu since, as stated above, epsilon had already been used to designate the chromium-rich body-centered cubic phase. The hitherto unknown ternary phases discovered in the 1200° C isothermal sections of the chromium-cobalt-molybdenum and the chromium-nickel-molybdenum ternary systems have been named R and P, respectively.

This work was conducted at the University of Notre Dame under the sponsorship and with the financial assistance of the National Advisory Committee for Aeronautics. The authors wish to thank Mr. Francis Pall for doing some of the work relating to the chromium-cobalt-molybdenum and chromium-nickel-molybdenum systems. The assistance of Messrs. C. Patrick Sullivan and Robert Hochman is also appreciated.

LITERATURE SURVEY

The chromium-cobalt-nickel-molybdenum quaternary system comprises the following six binary and four ternary systems: Chromium-cobalt, chromium-nickel, cobalt-nickel, chromium-molybdenum, cobalt-molybdenum, nickel-molybdenum, chromium-cobalt-nickel, chromium-cobalt-molybdenum, chromium-nickel-molybdenum, and cobalt-nickel-molybdenum. All of the above binary systems were investigated previously. The chromium-cobalt system was investigated by Elsea, Westerman, and Manning (reference 3), and the most recent chromium-nickel phase diagram was reported by Jenkins, Bucknall, Austin, and Mellor (reference 4). The "Metals Handbook" (reference 5) gives the accepted cobalt-nickel diagram, which was thoroughly investigated by several workers.

The chromium-molybdenum binary system is also given in the "Metals Handbook" (reference 5). In recent work on molybdenum-rich alloys, Kessler and Hansen (reference 6) confirm the results of other investigators that chromium and molybdenum are completely soluble in the solid state. The cobalt-molybdenum binary system was investigated by Sykes and Graff (reference 7). The transformation from face-centered cubic to hexagonal in pure cobalt above 1000° C indicated in their diagram was not found by several later investigators (references 8 to 10). Henglein and Kohsok (reference 11) recently pointed out that the cobalt-molybdenum intermediate phase Co7Mo6 is isomorphous with the iron-molybdenum, cobalt-tungsten, and iron-tungsten intermediate phases Fe7Mo6, Co7W6, and Fe7W6. The crystal structure of the Co7Mo6

phase can be described as hexagonal or rhombohedral (reference 11). The nickel-molybdenum binary system, as determined by Ellinger, is given in reference 5.

Of the four ternary systems involved in the chromium-cobalt-nickel-molybdenum quaternery system, two were previously investigated. The 1200° C isothermal section of the chromium-cobalt-nickel ternary system was investigated by Manly and Beck (reference 1). This diagram was slightly modified in the chromium-cobalt sigma-phase-field region by Kamen and Beck (reference 2). Siedschlag (reference 12) investigated the chromium-nickel-molybdenum ternary system, but the results of that investigation proved to be of no help in the present work. The nickel-rich alloys were examined from a technological point of view only, and phase relationships were not studied in detail. It was, therefore, necessary to investigate the three ternary systems chromium-cobalt-molybdenum, chromium-nickel-molybdenum, and cobalt-nickel-molybdenum. The survey of these systems was confined to the determination of the boundaries of the face-centered cubic (alpha) solid solutions and of the phases coexisting with the alpha phase at 1200° C.

The brittle intermetallic sigma phase, which occurs in many systems involving the transition elements, has recently become the subject of much interest and investigation. Sully and Heal (reference 13) have pointed out that the iron-chromium sigma phase and the chromium-cobalt gamma phase are isomorphous, and Goldschmidt (reference 14) found that the iron-molybdenum zeta phase is also isomorphous with the iron-chromium sigma phase. Beck and Manly (reference 15) investigated the chromiumcobalt-iron and the chromium-cobalt-nickel ternary systems. They proved that the iron-chromium sigma phase and the chromium-cobalt gamma phase form an uninterrupted series of solid solutions across the chromiumcobalt-iron ternary isothermal section at 800° C. They also found thatthe chromium-cobalt gamma phase extends deep into the chromium-cobaltnickel ternary system at 1200°C in a manner suggesting that nickel atoms and cobalt atoms substitute for each other in forming the sigma phase, while the chromium content of the phase remains essentially unchanged. On the basis of these results and of the existence of the sigma phase in the iron-vanadium system, Beck and Manly (reference 15) suggested that the sigma phase should also occur in the cobalt-vanadium and the nickel-vanadium systems. Indeed, this was confirmed by Duwez and Baen (reference 16) who also formally postulated the criterion that a face-centered and a body-centered cubic metal are required in an alloy system before the sigma phase will form. In the present investigation, special attention was given to the occurrence of the sigma phase, and further confirmation was obtained for the earlier observation that atoms of elements of like structure can substitute for each other in the formation of the sigma phase. A criterion for the formation of the sigma lattice in terms of electron vacancy concentration in the 3d subband was derived in order to rationalize the above observations.

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EXPERIMENTAL PROCEDURE

In this work, the phase boundaries were established by microscopic examination of carefully homogenized alloys. The phases were identified by means of X-ray diffraction. Details relating to the equipment used and to the melting and homogenizing procedures followed throughout the work were reported by Manly and Beck (reference 1). Molybdenum in the form of a 1/8-inch-diameter rod was used to make up the alloys. The lot analyses of the electrolytic chromium, of the cobalt rondelles, and of the nickel used are given in table I.

Most of the alloys were melted in Alundum crucibles, except for 34 alloys melted in zirconia and stabilized zirconia crucibles at the beginning of the work. The type of crucible used to melt each alloy is listed in table II. Early in the work, it was found that molybdenumbearing alloys were susceptible to zirconium pickup. This was confirmed by semiquantitative spectrographic analysis. The use of zirconia and stabilized zirconia crucibles was, therefore, discontinued, and all subsequent alloys were melted in Alundum crucibles. The zirconium pickup is treated in greater detail under Discussion.

The ingots were generally found to be free of excessive segregation. except in a few isolated cases discussed later. Specimens for homogenization and subsequent microscopic and X-ray analysis were taken from the bottom section of each ingot. Immediately adjacent examples were used for chemical analysis. All specimens which consisted mainly of the face-centered cubic (alpha) solid solution were double-forged prior to homogenization. This double-forging treatment consisted of heating the specimen at 1200° C for 1/2 hour, forging, reheating at 1200° C for 1/2 hour, and forging again. With this preliminary treatment alpha alloys were easily homogenized by annealing at 1200° C for 48 hours. Alpha alloys containing more than about 20 percent of any second phase were too brittle to be forged. Specimens from such alloys were homogenized for 95 to 150 hours at 1200° C. All specimens were quenched directly into cold tap water. It is very important that the oxygen be removed from the furnace atmosphere in which alloys containing molybdenum are being annealed, because of the extremely rapid rate of oxidation of these alloys at high temperatures. It is believed that the homogenizing treatments used gave very nearly equilibrium conditions, except in a few cases to be discussed later, because continued annealing, which in some cases was extended up to 200 hours, resulted in no detectable further changes in the microstructures.

After homogenization, a powder for X-ray analysis was taken from each specimen, by either filing or crushing, depending on the brittleness of the alloy, and the remaining piece was prepared for microscopic examination to detect the presence or absence of a second phase. The phase boundaries were thus determined by the disappearing phase method.

It was found necessary to vary the etching procedure considerably, according to the composition of the alloy.

(1) The following etchants were used successfully to differentiate between the various phases:

Hydrochloric acid, milliliters		•		•		•	•	•	•	•	•		8.0
Nitric acid, milliliter													0.5
Glycerin, milliliters													2.0
Cupric chloride, milligrams										5	50	to	150

The freshly polished specimen was either immersed in the etchant for several minutes, or, for more rapid attack, swabbed with cotton saturated with the etchant. This etchant was used only to reveal the structure of the alpha phase; grain boundaries, annealing twins, and transformation striations became apparent, and any second-phase particles present were clearly delineated. It was not possible, however, to identify which second phase was present by using this etchant alone.

(2) The following electrolytic etching and staining procedure was used with alloys in the chromium-nickel-molybdenum ternary system:

Oxalic acid, grams	8
Distilled water, milliliters	
Cathode	Stainless steel
Voltage, volts (d-c.)	6
Electrode spacing, inch	
Temperature, OC	
Time, seconds	

The specimen was removed from the etching bath and immediately immersed for 10 to 20 seconds into a staining solution consisting of 5 grams of potassium permanganate and 5 grams of sodium hydroxide dissolved in 90 milliliters of distilled water. This etching and staining method worked well with the epsilon phase. Grain boundaries became evident and the characteristic Widmanstätten precipitate of sigma in the epsilon phase, when present, became clearly recognizable. The epsilon phase was stained a very light tan color, or a darker brown when the Widmanstätten precipitate was present. The alpha phase always remained unattacked and unstained. The chromium-nickel-molybdenum ternary sigma phase always stained, and a faint structure was sometimes brought out in this phase. The color of the stain on the sigma phase varied from bright green or red to purple. The grain boundaries of the ternary P phase were attacked and this phase, too, was always stained, the color varying from green to red and orange. The delta phase was generally very unevenly attacked and either did not stain or stained an uneven brownish color. Grain boundaries were revealed in the delta phase. With chromium-nickel-molybdenum alloys, great success was experienced in differentiating between epsilon, sigma, and alpha when

these phases coexisted. As a result of this, it was possible to locate the corner of the three-phase alpha-plus-epsilon-plus-sigma field on the alpha boundary by metallographic means. Even when present as small particles in an alpha matrix, sigma consistently stained a brilliant color, whereas the epsilon phase never stained more than a light tan color. It was difficult to distinguish between minor amounts of the sigma and the P phases in an alpha matrix because of their similar staining characteristics. However, when sigma and P were adjacent in the same alloy, it was possible to distinguish one phase from the other. Final identification, of course, was made by means of X-ray analysis.

(3) The electrolytic oxalic-acid etchant described above proved unsatisfactory for chromium-cobalt-molybdenum alloys. The following electrolytic etchant, used in conjunction with the previously described staining solution, was used with satisfactory results:

Concentrated phosphoric acid,	milliliters		, 5
Distilled water, milliliters			
Cathode			
Voltage, volts (d-c.)			. 6
Electrode spacing, inch			
Temperature, OC		20 to	30
Time, seconds		10 to	15

The alpha phase was lightly attacked but remained unstained. The sigma phase was lightly attacked and stained colors varying from orange to blue or purple. A fine Widmanstätten precipitate was sometimes observed in the sigma phase. The chromium-cobalt-molybdenum ternary R phase was unattacked, but this phase stained yellow to rusty brown. The mu phase was attacked and usually stained pale blue or did not stain at all. Grain boundaries and tiny annealing twins were observed in the mu phase. Sigma, R, and alpha were easily distinguished from each other when these phases occurred together in the same alloy, but small particles of sigma were indistinguishable from small particles of either the R or the mu phase in an alpha matrix. The mu phase was readily differentiated from R.

(4) It was necessary to decrease the acidity of the phosphoric-acid electrolyte and increase the alkalinity of the alkaline permanganate staining solution in order to differentiate successfully between the delta and the mu phases in cobalt-nickel-molybdenum alloys. The following solution was used:

Concentrat	ed phos	sphoric	acid,	mi.	1113	.it€	ers	.•	•	•	•	•		•	•	•	. 2	2.5
Distilled '	water.	millili	Lters		٠.												9"	7.5

Etching conditions were the same as those for procedure 3. The specimen was then immersed into a staining solution containing 20 grams of sodium

hydroxide and 5 grams of potassium permanganate dissolved in 75 milliliters of distilled water. The alpha phase was lightly attacked but remained unstained. The delta phase was unevenly attacked, grain boundaries being revealed, and was stained dark green or blue. The mu phase was more evenly attacked; grain boundaries and occasional twins were evident. The mu phase stained a number of different colors, apparently depending upon the orientation, each grain or twin showing only one color. With this etching procedure it was possible to detect small amounts of the mu phase in a matrix of delta but, because of the wide variation of color from grain to grain in the mu phase, it was not possible to identify small amounts of delta in a matrix of the mu phase. This difficulty of microscopically identifying small amounts of the delta phase in a matrix of mu was circumvented by the use of an X-ray method to determine the solubility limit of nickel in the mu phase. In a plot of the d values for one high-angle diffraction line in a series of mu alloys as a function of the nickel content, the solubility limit is indicated where the slope of the curve becomes zero.

The powders for X-ray analysis were sealed under vacuum in fusedquartz capsules and annealed at 1200° C for 1/2 hour. The capsules were then quenched directly into cold tap water. The powders were mounted on 1- by 1-inch cards with a colloidal glue, and a diffraction pattern was taken in an asymmetrical focusing camera of 20-centimeter diameter, using unfiltered chromium radiation at 8 milliamperes and 30 kilovolts. Thus, the identity of the phases present in each homogenized alloy was confirmed by X-ray diffraction. However, the X-ray method of identification was found relatively insensitive to small amounts of a second phase. especially in alloys having an alpha matrix. This difficulty was surmounted partially by careful preparation of the powder. A large quantity of filings was collected from the specimen to insure obtaining a representative sample. The minor phase was generally much more brittle than the alpha matrix and had a much smaller particle size. It was concentrated by sieving the powder to 200 mesh. Only the finest powder from the sample was used to prepare the X-ray specimen. In handling this fine powder precautions were necessary in order to avoid losing any of the minor phase. For example, the original filings had to be collected on a smooth tracing paper or Celluloid, to which the powder will notadhere and from which the fine powder may be easily recovered.

An X-ray method of locating three-phase-field corners on the alpha phase boundary, as described previously by Manly and Beck (reference 1), was used in this work to supplement the microscopic results. The method consists of plotting the lattice parameters of the saturated alpha alloys as a function of composition. The three-phase-field corner is usually indicated by a rather sharp change in the slope of the curve. The lattice parameters for these plots were determined by taking back-reflection diffraction patterns from the surface of microspecimens known to be saturated alpha. Flat film and a collimating system of 1-millimeter diameter were

used. By using unfiltered chromium radiation and a specimen-to-film distance of 29.54 millimeters, the chromium K α reflections from the (220) planes and the chromium K β reflections from the (311) planes of the alpha phase were recorded. Smooth circles were obtained by rotating the specimen about an axis parallel to that of the collimating system but offset from it. The lattice parameter determined from each of these lines was plotted against $\cos 2\theta (2\cos^2\theta - 1)$ and extrapolated to $\theta = 90^\circ$ in order to eliminate systematic errors.

EXPERIMENTAL RESULTS

The experimental data for all alloys investigated are given in tables III to VI. The 1200°C isothermal sections for the cobalt-nickel-molybdenum, chromium-cobalt-molybdenum, and chromium-nickel-molybdenum ternary systems and the solubility limits of the alpha phase in the 2.5-, 5-, 10-, and 20-percent-molybdenum quaternary sections, as shown in figures 1 to 8, were drawn in accordance with these data. The amount of each phase, as estimated microscopically in the homogenized structure, is tabulated together with corresponding X-ray diffraction data for each alloy. The amounts of the phases, corresponding to the phase diagrams, are also given for comparison. For alloys which were chemically analyzed the content in acid insoluble material was determined and found to vary from nil to a few tenths of 1 percent. The compositions of chemically analyzed alloys reported in the tables were corrected to 100-percent-metal content.

Phases

The various phases identified in this investigation and the metallographic characteristics of each phase are described in the following paragraphs.

Alpha. The alpha phase, which forms the matrix of most of the practically important high-temperature alloys, is based on solid solutions of the face-centered cubic elements cobalt and nickel. This phase, being relatively soft and ductile, was easily hot-forged, and alpha alloys rich in nickel could be cold-worked to some extent. Reagent 1 was used to reveal the microstructure of the alpha phase. The structure is typical of most face-centered cubic solid solutions, showing equiaxed grains and numerous annealing twins. In alpha alloys containing small amounts of a second phase a great grain-size contrast was sometimes seen. Such "duplex" structures undoubtedly occur as a result of the inhibition effect of small second-phase particles on grain growth (reference 17). On cooling, pure cobalt is known to transform from the face-centered cubic to the hexagonal close-packed structure by a martensitic transformation (reference 8). In cobalt-rich alpha solid solutions the beginning of this transformation is manifested in the microstructure by the presence of

striations, as illustrated in figure 9. These transformation striations were also observed in saturated cobalt-molybdenum and cobalt-chromium alpha alloys and in quaternary alpha alloys rich in cobalt. Additions beyond about 15 percent nickel to cobalt-molybdenum and 30 percent nickel to cobalt-chromium saturated alpha alloys seem to suppress the transformation, as evidenced by the absence of striations in the microstructure.

Microspecimens of alloys containing alpha in addition to any other phases exhibited a pronounced relief effect in the as-polished and unetched condition. The relief is due to the large difference in hardness between alpha and the phases which coexist with it at 12000 C. unetched microspecimens, the alpha phase had a slightly amber or yellowish tint, while the other phases appeared white. Figure 10 shows a banded structure which was observed in alloys consisting of approximately equal amounts of the alpha and mu phases. Figure 11 illustrates the general shape and distribution of minor amounts of alpha in a matrix of the sigma phase. This is typical of the distribution of minor amounts of alpha in all phases. Alloys which consisted of larger amounts of alpha coexisting with any of the other phases exhibited microstructures similar to the one shown in figure 12. Data from a typical X-ray diffraction pattern of the alpha phase are given in table VII. Evidence that preferred orientation may occur in forged and annealed alpha alloys was found in some of the back-reflection pictures taken from the surface of microspecimens.

Epsilon. - The body-centered cubic epsilon phase at 1200° C is based on solid solutions of chromium and molybdenum. Within the composition ranges investigated in this work, the epsilon phase was encountered only in the chromium-nickel-molybdenum ternary isothermal section where it coexists with the alpha and sigma phases and in quaternary alloys containing less than about 3 percent molybdenum near the chromium-nickel side of the diagram, where it coexists with alpha. When etched and stained according to procedure 2 the microstructure of the epsilon phase was clearly revealed. This phase stained a light tan color and, in this work, a characteristic Widmanstätten type of precipitate was always observed in the large equiaxed grains of the epsilon phase, with the exception of epsilon coexisting with alpha in alloys near the chromiumnickel binary side of the diagram. This Widmanstätten precipitate was previously found in chromium-rich alloys in the epsilon phase fields of both the chromium-cobalt-nickel and the chromium-cobalt-iron ternary isothermal sections at 1200°C (references 1 and 2) and was identified in both cases as being the sigma phase. In the present work, too, it has been concluded that the Widmanstätten precipitate consists of the sigma phase. In epsilon alloys very near to the corner of the threephase alpha-epsilon-sigma field in the chromium-nickel-molybdenum ternary isothermal section, the Widmanstatten precipitate in the epsilon phase is extremely heavy. (See fig. 13.) The epsilon phase was easily distinguished from alpha or sigma, either by the precipitate in epsilon or by its tan color, when stained (procedure 2), since the alpha phase remains unstained and the sigma phase stains to a brilliant color. Figure 14 is an example of this, where the three phases are seen to coexist.

This alloy was slightly deformed prior to microscopic examination. Note the severe cracks in the heavily stained sigma and the few cracks in the lightly stained epsilon. Table VIII gives the data from a typical X-ray diffraction pattern of the epsilon phase.

Sigma. The sigma phase is an extremely brittle intermetallic phase which, in this investigation, occurred in the 1200° C isothermal sections of the chromium-cobalt-molybdenum and chromium-nickel-molybdenum ternary systems; as well as in chromium-cobalt-nickel-molybdenum quaternary alloys containing up to 20 percent molybdenum. The sigma phases in these, as in other systems, are isomorphous. Recent investigations on single crystals of the chromium-cobalt and iron-chromium sigma phases have yielded information showing that the crystal class of sigma is tetragonal. 1 Microspecimens from sigma alloys invariably showed cracks as a result of the brittleness of the phase. Qualitatively, ingots of chromium-nickel-molybdenum ternary sigma alloys did not seem to be quite so brittle as chromium-cobalt-molybdenum sigma alloys in that they did not shatter quite so easily. Ingots of chromium-cobalt-molybdenum sigma alloys were so brittle that they shattered when dropped to the floor. Etching and staining the chromium-nickel-molybdenum ternary sigms phase was accomplished by following procedure 2. Prolonged etching brought out faintly the grain boundaries in sigma, as shown in figure 15. It is interesting to note that, in specimens which consisted almost entirely of the sigma phase, the color of the stain on sigma was very light, whereas massive particles of sigma in a matrix of some other phase always stained a brilliant color. The reason for this is not clearly understood. In the two-phase sigma-plus-P field of the chromiumnickel-molybdenum system at 1200° C, the grains of these two phases were very intimately arranged so that grain boundaries seemed to correspond to phase boundaries. This is illustrated in figure 16. The microstructure of chromium-cobalt-molybdenum alloys in which the sigma phase coexisted with minor amounts of the alpha or R phases was best revealed by etching and staining, as outlined in procedure 3. The grain boundaries of the sigma phase were not attacked, but a scattered Widmanstätten precipitate was sometimes brought out. (See fig. 17.) The identity of this precipitate is not known.

Sigma coexists with the alpha, epsilon, and R phases in randomly distributed particles and with the epsilon phase in some cases as a Widmanstatten precipitate. Small particles of the sigma phase which precipitated along the grain boundaries from a supersaturated chromium-cobalt-molybdenum alpha alloy are shown in figure 18. Etching procedure 3 was used; consequently, the structure of the alpha was not revealed. Data from typical X-ray diffraction patterns of the chromium-nickel-molybdenum and chromium-cobalt-molybdenum sigma phases are listed in table IX. It is seen from a comparison of the two patterns that these sigma phases are isomorphous.

Reference 18; also private communication from Messrs. J. S. Kasper, B. F. Decker, and J. R. Belanger, General Electric Research Laboratory.

naca tn 2683

Delta.- Solid solutions based on the nickel-molybdenum delta phase are formed in the 1200° C isothermal sections of the chromium-nickelmolybdenum and cobalt-nickel-molybdenum ternary systems. The crystal structure of the delta phase is not known. This phase is hard and brittle, although not nearly so brittle as the sigms phase. Ingots of alloys consisting mainly of the delta phase could not be broken by a sharp blow with a hammer. Fine cracks were sometimes seen in the microstructure, which was revealed by etching procedure 4. These cracks probably originated during the preliminary grinding of the microspecimen. Figure 19 is a photomicrograph of an alloy of delta with minor amounts of alpha and mu. This specimen was etched and stained according to procedure 4. Note the large grains of delta and the much smaller grains of mu which are stained to various colors and exhibit occasional tiny annealing twins. The alpha particles are slightly roughened but remained unstained and appear white. As the amount of the mu phase coexisting with delta increases, it becomes increasingly difficult to distinguish between the two phases. In chromium-nickel-molybdenum alloys consisting largely of delta, no difficulty was experienced in differentiating between the delta and P phases. Small particles of the P phase in a matrix of delta are shown in figure 20. Data from a typical X-ray diffraction pattern of the delta phase are listed in table X.

Mu. - In the present work, solid solutions based on the intermediate cobalt-molybdenum epsilon phase were renamed mu to avoid contradictions in the nomenclature. This phase occurs in the 1200° C isothermal sections of the chromium-cobalt-molybdenum and cobalt-nickel-molybdenum ternary systems. The mu phase also coexists with alpha in chromium-cobaltnickel-molybdenum quaternary alloys from at least 17 to 20 percent molybdenum near the cobalt-nickel side of the diagram. The crystal structure of the mu phase can be described as hexagonal or rhombohedral (reference 11). This phase is isomorphous with the corresponding intermediate phase in the iron-molybdenum, iron-tungsten, and cobalt-tungsten binary systems (reference 11). Ingots of mu alloys displayed physical characteristics similar to those of ingots of delta alloys. A macrostructure was generally visible, without etching, on the surface of the ingots. The macrostructure was either granular or dendritic in nature. No relief effect was observed in as-polished and unetched microspecimens containing minor amounts of the delta or R phases in a matrix of mu, indicating that these phases have nearly the same hardness. In attempting to reveal clearly the microstructure of the mu phase it was found that large areas throughout the specimen were preferentially attacked, giving one the impression that the alloy contained two phases when actually there was only one. (This was proved repeatedly by careful X-ray diffraction studies.) After experimenting with various etching techniques, all with the same result, it was concluded that the preferential attack is an orientation effect. Figure 21 shows the structure of a typical mu alloy. This specimen was etched according to procedure 4 but omitting the stain.

13

The mu phase coexists with the delta and R phases in randomly distributed particles. A banded structure in alloys of mu plus alpha has been discussed and illustrated previously in figure 10. Table XI gives the data from a typical X-ray diffraction pattern of the mu phase.

P phase. - The P phase was discovered in the 1200° C isothermal section of the chromium-nickel-molybdenum ternary system. This phase is not known to occur in any of the three binary systems at any temperature. The crystal structure of the P phase was not determined. Alloys of P were hard and brittle, having much the same physical characteristics as alloys of the chromium-nickel-molybdenum ternary sigma phase. Numerous cracks were observed in the microstructure, which was very similar to that of the ternary sigma phase. Grain boundaries in the P phase were attacked when etched by procedure 2, and this etchant also produced stain colors varying from green to red and orange. Because of the similarity in microstructures and staining characteristics of the P and sigma phases, it was difficult to distinguish between them microscopically. In figure 22, minor amounts of sigma and P are seen in an alpha matrix. Sigma is stained darker than P, but the contrast is slight. Small particles of the delta or alpha phases could easily be identified in a matrix of the P phase. Data from a typical X-ray diffraction pattern of the P phase are given in table XII.

R phase. The R phase was discovered in the 1200°C isothermal section of the chromium-cobalt-molybdenum ternary system. The crystal structure of the R phase was not determined. Ingots of alloys which consisted of the R phase had about the same physical characteristics as delta and mu alloy ingots. When etched and stained according to procedure 3, the R phase was unattacked but stained yellow to rusty brown. Small quantities of the sigma, mu, or alpha phases in a matrix of R were easily identified microscopically. Figure 23 shows alpha and a few particles of the sigma phase in a matrix of the R phase. The sigma particles were stained but not so clearly delineated as unstained alpha particles. The contrast achieved by procedure 3 is further illustrated in figure 24, where larger amounts of alpha and sigma coexist in a matrix of the R phase. Data from a typical X-ray diffraction pattern of the R phase are presented in table XIII.

Impurity phases.— Three main impurity phases were observed in the microstructures of alloys investigated in this work. Metallic-oxide inclusions (mostly chromium oxide) were present in the alloys in varying amounts. Chromium-rich epsilon alloys showed the highest number of oxide inclusions, while alloys rich in nickel and molybdenum showed the least. Metallic-oxide inclusions are easily recognized in as-polished and unetched microstructures. They are generally unattacked by etchants. Chromium-oxide particles are visible in figure 13 which is an epsilon alloy etched to bring out the Widmanstätten precipitate of sigma.

Another impurity phase appeared in some alloys which were melted in zirconia or stabilized zirconia crucibles. The amount of this impurity phase was observed to be especially high in an alloy which had been slightly overheated during melting in a stabilized zirconia crucible. The specimen had been forged and homogenized at 1200° C for 48 hours. The microstructure showed small amounts of sigma second phase associated with a dark etching phase in a network outlining the grains of alpha, suggesting a eutectic origin. This structure is illustrated in figure 25. This alloy was analyzed by quantitative spectrographic analysis and was found to contain more than 1.00 percent zirconium. It was concluded that the impurity phase resulted from the reaction between the molybdenum-containing liquid metal and the crucible.

Minute amounts of a bright yellow impurity phase were found in all alloys which were melted in zirconia or stabilized zirconia crucibles. It is believed that this impurity, too, resulted from zirconium pickup by the alloy. Spectrographic analysis of an alloy melted in an Alundum crucible revealed no aluminum pickup. Microscopic observation did not show any impurity connected with aluminum in any alloy melted in an Alundum crucible. All alloys in the latter part of this work were melted in Alundum crucibles.

Phase Diagrams

The various phase diagrams determined in the present work are described in the following paragraphs.

Cobalt-nickel-molybdenum ternary system at 1200° C.- The 1200° C isothermal section of the cobalt-nickel-molybdenum ternary system, as drawn from alloy data listed in table III, is presented in figure 1. For convenience, the same diagram with the alloy compositions indicated is shown in figure 2. Within the composition ranges investigated, only solid solutions of the phases known from the binary systems were found; no ternary phases occur. About two-thirds of the diagram is covered by the alpha phase field and the two-phase alpha-plus-mu field. The mu phase field penetrates deep into the ternary isothermal section, roughly parallel to the cobalt-nickel side. Limited solid solutions of delta coexist with the mu and alpha phases in two narrow two-phase fields. Alpha, delta, and mu coexist in a three-phase field pointing toward the nickel corner of the diagram.

The face-centered cubic solid solutions based on the cobalt-nickel slpha phase extend to 36 percent molybdenum on the nickel-molybdenum side and to 22.5 percent molybdenum on the cobalt-molybdenum side of the diagram. This is in good agreement with the published nickel-molybdenum and cobalt-molybdenum binary systems. The solubility limit of the alpha phase in the ternary isothermal section is slightly concave, running from the cobalt-molybdenum side to the corner of the three-phase

alpha-delta-mu field (point 10, fig. 1), and almost a straight line from the corner of the three-phase field to the nickel-molybdenum binary diagram. The corner of the three-phase field on the alpha boundary (point 10, fig. 1) is placed at 10 percent cobalt, 35 percent molybdenum, and 55 percent nickel. This point was located by combined microscopic and X-ray data. Alloy 627 contained very small amounts of both the delta and mu phases. In order to check the microscopic findings, the lattice parameters of saturated alpha alloys along the boundary were plotted as a function of cobalt content. The corner of the three-phase field is usually indicated by a sudden change in the slope of the parameter curve. This plot is presented in figure 26 from data listed in table XIV. It is evident from figure 26 that the general effect of cobalt going into solid solution in the saturated nickel-molybdenum alpha phase is to contract the lattice. Unfortunately, however, there were not enough points on the parameter curve to locate the cusp accurately. The point at which the parameter curve suddenly changes slope was definitely located by determining the parameter of the alpha which coexists in the three-phase field. Since the composition of the alpha is constant anywhere in the three-phase field, the lattice parameter is also constant. The composition corresponding to this point on the parameter curve (point 10, fig. 26) gives the location of the threephase-field corner. Within the limits of experimental accuracy, this point lies between 9.50 and 10.50 percent cobalt. This result is in excellent agreement with the microscopic findings. Tests were made on the reproducibility of parameter measurements on the same alloy. The lattice parameter of an alpha alloy which gave diffraction rings that were particularly difficult to read (intensity maximums due to preferred orientation) was determined from five different films, each one taken after replacing the specimen on the camera. The maximum variation in these five measurements was 0.001 kX unit. A similar check on an alloy which produced sharp, smooth diffraction rings gave a maximum variation in five measurements of only 0.0003 kX unit.

In alloy 407 a large discrepancy was found between the intended melted analysis and the reported chemical analysis. Both analyses are reported in table III. The reported chemical analysis, however, moves the location of the alloy almost parallel to the alpha boundary so that the course of the boundary is not changed. The intended analysis is plotted in figure 2.

Solid solutions based on the mu phase extend to at least 24 percent nickel (point 11, fig. 1) in the cobalt-nickel-molybdenum 1200° C isothermal ternary section. This was confirmed by X-ray investigation. As nickel goes into solid solution, the lattice of the mu phase expands gradually, becoming constant in the two-phase delta-plus-mu field and in the three-phase field. The d values for a high-angle X-ray line in a series of mu alloys were plotted as a function of nickel content, as shown in figure 27. The solubility limit is indicated at 24 percent nickel (point 11, fig. 27). The d values for the same line in patterns

16 - NACA TN 2683

of mu for alloys located in the three-phase field were determined as a further check. As seen from the data given in table XV, the agreement was good.

The alpha and mu phases coexist in a large two-phase field extending from the cobalt-molybdenum side of the diagram to the corners of the three-phase field on the alpha and mu phase boundaries. The corner of the three-phase field on the mu phase boundary (point 11, fig. 1) is placed at 17 percent cobalt, 24 percent nickel, and 59 percent molybdenum.

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The two-phase mu-plus-delta field is a narrow field between the ends, of the mu and delta phases.

The delta phase extends to only 6.5 percent cobalt in the isothermal section and coexists with alpha in a two-phase field which extends from the nickel-molybdenum side of the diagram to the corners of the three-phase field on the alpha and delta phase boundaries. The corner of the three-phase field on the delta phase boundary (point 12, fig. 1) was found to be at 6.5 percent cobalt, 33.5 percent nickel, and 60 percent molybdenum.

The composition range of the three-phase alpha-delta-mu field is outlined by the three corners (points 10, 11, and 12, fig. 1) given above. The obtuse angle of this triangular field is at the delta corner.

Chromium-cobalt-molybdenum ternary system at 1200° C.- Figure 3 is the 1200° C isothermal section of the chromium-cobalt-molybdenum ternary system, drawn in accordance with the alloy data listed in table_IV. The same diagram is presented in figure 4 with alloy compositions indicated. In addition to solid solutions of the phases known from the binary systems, a new ternary phase, not known in any of the binary systems. was discovered. This new ternary phase was designated as the R phase. Solid solutions of the face-centered cubic alpha phase surround the cobalt corner of the diagram. The chromium-cobalt sigma phase extends approximately halfway across the isothermal section in a field roughly parallel to the chromium-molybdenum side of the diagram, and solid solutions based on the cobalt-molybdenum mu phase occur in a small field, also approximately parallel to the chromium-molybdenum side. The R phase field is located between the ends of the sigma and mu phase fields. The extension of the mu and R phase fields in the direction parallel to the cobalt-molybdenum side of the diagram was not investigated in detail. Alpha coexists with the sigma and mu phases in two wide two-phase fields and with the R phase in a long, narrow two-phase field. The R phase coexists with the sigma and mu phases in two narrow two-phase fields. The alpha, sigma, and R phases coexist in a threephase field between the alpha-sigma and alpha-R two-phase fields. Alpha, R, and mu coexist in a three-phase field between the two-phase fields of alpha and R and of alpha and mu.

The solubility of chromium in cobalt at 1200° C was determined by Manly and Beck (reference 1) at 34 percent. In the present work, the solubility of molybdenum in cobalt was found to be 22.5 percent. This value is approximately 1.00 percent lower than that indicated by the cobalt-molybdenum diagram reported by Sykes and Graff (reference 7). The difference is of the order of magnitude of experimental error. The alpha phase boundary in the 1200° C. isothermal ternary section is a convex curve from the chromium-cobalt binary to the corner of the threephase alpha-sigma-R field (point 13, fig. 3) and almost a straight line from the cobalt-molybdenum binary to the corner of the alpha-sigma-R three-phase field. There is a slight cusp on the alpha boundary at the corner of the three-phase alpha-R-mu field (point 16, fig. 3). The alpha-sigma-R three-phase-field corner (point 13, fig. 3) is placed at 21.0 percent chromium, 66 percent cobalt, and 13 percent molybdenum, while the corner of the alpha-R-mu three-phase field (point 16, fig. 3) is at 16 percent chromium, 69 percent cobalt, and 15 percent molybdenum. In figure 28, the lattice parameters of saturated alpha alloys along the alpha phase boundary are plotted as a function of chromium content. Data for this plot are given in table XVI. The three-phase-field corners on the alpha phase boundary are clearly indicated by sharp breaks in the parameter curve. The breaks occur at approximately 14.5 and 21.0 percent chromium (points 13 and 16, fig. 28) which agrees well with the microscopic work. (The reason for the sudden decrease and increase in parameters at the corners is explained under Discussion.)

The sigma phase field is a narrow band, from 40 to 45 percent cobalt, which penetrates to a maximum of 31 percent molybdenum in the 1200° C isothermal ternary section. The corner of the three-phase alpha-sigma-R field on the sigma phase boundary (point 14, fig. 3) is at 25 percent chromium, 46.5 percent cobalt, and 28.5 percent molybdenum. Evidence was found in the microstructure of alloy 531 that another three-phase-field corner occurs on the sigma phase boundary at approximately 27 percent chromium, 42 percent cobalt, and 31 percent molybdenum. This may be a corner of a three-phase sigma-epsilon-R field. It was observed from X-ray diffraction patterns of the sigma phase that the lattice expands as the amount of molybdenum in solid solution increases. No quantitative plot of d values against composition was made to check the solubility limit, however, because the end of the sigma phase field was easily located by microscopic means.

The two-phase alpha-plus-sigma field extends from the chromium-cobalt side of the diagram to the corners of the three-phase alpha-sigma-R field on the alpha and sigma phase boundaries.

The third corner of the alpha-sigma-R three-phase field (point 15, fig. 3) is on the boundary of the R phase at 18 percent chromium, 44 percent cobalt, and 38 percent molybdenum. The obtuse angle in this triangular field is at the sigma corner.

naca in 2683

The solubility limits of the R phase in the direction parallel to the chromium-molybdenum side of the diagram were located at approximately 14.5 and 19.5 percent chromium. The alpha-R-mu three-phase-field corner on the R boundary (point 17, fig. 3) is placed at 13.5 percent chromium, 44 percent cobalt, and 42.5 percent molybdenum. The other corner of this very narrow three-phase field (point 18, fig. 3) is on the boundary of the mu phase at 9.5 percent chromium, 42.5 percent cobalt, and 48 percent molybdenum. The obtuse angle in this triangular field is at the corner on the R phase boundary.

The alpha-plus-R two-phase field is located between the three-phase fields of alpha, sigma, and R and of alpha, R, and mu.

The mu phase takes a maximum of approximately 11 percent chromium into solid solution and coexists with alpha in a large two-phase field extending from the cobalt-molybdenum binary to the corners of the three-phase alpha-R-mu field on the alpha and mu phase boundaries.

Chromium-nickel-molybdenum ternary system at 1200° C.- The 1200° C isothermal section of the chromium-nickel-molybdenum ternary system was drawn from data given in table V and is presented in figure 5. The same diagram with the alloy compositions indicated is shown in figure 6. Beside the phases known from the binary systems two new ternary phases were found to occur in the composition ranges investigated. One of these ternary phases has the familiar sigma structure and was, therefore, designated sigma. The crystal structure of the other new ternary phase, designated as the P phase, is unknown. About three-quarters of the area of the diagram, within the range of compositions investigated, is covered by solid solutions of the alpha phase around the nickel corner and by two-phase fields and three-phase fields in which alpha coexists with the epsilon, sigma, P, and delta phases. The large two-phase fields of alpha and sigma and of alpha and P are separated by a very narrow three-phase alpha-sigma-P field. A narrow three-phase alpha-Pdelta field lies between the alpha-P and alpha-delta two-phase fields. The alpha and epsilon phases coexist in a two-phase field along the chromium-nickel side of the diagram. The three-phase alpha-sigma-epsilon field, which separates the alpha-epsilon and alpha-sigma two-phase fields, is relatively large in comparison with the other three-phase fields. The ternary sigma phase coexists with epsilon and P in 2_two-phase fields. Delta and P coexist in a small two-phase field.

At 1200° C, solid solutions of face-centered cubic nickel in the chromium-nickel binary system extend to 57.5 percent chromium and in the nickel-molybdenum binary system, to 36 percent molybdenum. The alpha phase boundary in the ternary isothermal section has a sharp peak at the alpha corner of the three-phase alpha-epsilon-sigma field (point 19, fig. 5). This point was located by microscopic means at 43 percent chromium, 53.5 percent nickel, and 3.5 percent molybdenum. Except for a slight cusp at the alpha corner of the other three-phase

fields (points 22 and 25, fig. 5), the alpha boundary is fairly straight from the nickel-molybdenum binary to the alpha corner of the three-phase alpha-epsilon-sigma field. From data listed in table XVII, the lattice parameters of saturated alpha alloys along the alpha boundary were plotted in figure 29 as a function of chromium content. The alphaepsilon-sigma three-phase-field corner on the alpha boundary (point 19, fig. 5) is very clearly indicated at about 43 percent chromium by the drastic break in the curve (point 19, fig. 29). The curve is almost vertical beyond this point because the chromium content of saturated alpha coexisting with the epsilon phase varies between only 43 and 42.5 percent. The break in the parameter curve at 23 percent chromium (point 22, fig. 29), though not so drastic, is also in good agreement with the location of the three-phase alpha-sigma-P field corner (point 22, fig. 5) from microscopic findings. The alpha corner of the three-phase alpha-P-delta field (point 25, fig. 5) is not manifested by a pronounced cusp and is almost imperceptible. This point was located at 6 percent chromium, 61.5 percent nickel, and 32.5 percent molybdenum by carefully determining the boundaries of the three-phase alpha-P-delta field. Extrapolated segments of the parameter curve indicate a break at approximately 7.5 percent chromium (point 25, fig. 29), in fair agreement with the above observations.

The epsilon boundary was determined only along the two-phase alphaepsilon field. This boundary extends from the chromium-nickel side of the diagram to the epsilon corner of the three-phase alpha-epsilon-sigma field (point 20, fig. 5). As shown in figure 5, alpha, epsilon, and sigma coexist in a three-phase field whose boundaries are as follows: The alpha corner (point 19, fig. 5) at 43 percent chromium, 53.5 percent nickel, and 3.5 percent molybdenum; the sigma corner (point 21, fig. 5) at 49 percent chromium, 28 percent nickel, and 23 percent molybdenum; and the epsilon corner (point 20, fig. 5) at 60 percent chromium, 29.5 percent nickel, and 10.5 percent molybdenum.

The sigma phase field is approximately parallel to the chromium-molybdenum side of the diagram, comprising a range of about 23 to 49 percent chromium. As molybdenum replaces chromium in the sigma structure, the lattice expands. This is clearly illustrated in figure 30, which was plotted from the data given in table XVIII. The variation of one high-angle d value for a series of saturated sigma alloys along both sides of the sigma field is shown as a function of chromium content. The two curves meet at approximately 48 percent chromium (point 21, fig. 30), corresponding closely to the sigma corner of the three-phase alpha-epsilon-sigma field (point 21, fig. 5).

Solid solutions of the new ternary P phase exist between 8 and 19 percent chromium. The two-phase fields of P and sigma and of P and delta are very narrow, each one extending over a range of only about 4 percent chromium.

The corners of the three-phase alpha-sigma-P field are located at the following points: The alpha corner (point 22, fig. 5) at 23 percent chromium, 57.5 percent nickel, and 19.5 percent molybdenum; the sigma corner (point 23, fig. 5) at 22 percent chromium, 31 percent nickel, and 47 percent molybdenum; and the P corner (point 24, fig. 5) at 19 percent chromium, 32.5 percent nickel, and 48.5 percent molybdenum.

At 1200°C, the nickel-molybdenum delta phase dissolves only about 4 percent chromium. This phase coexists with alpha in a long, narrow, almost rectangular two-phase field along the nickel-molybdenum binary system.

The boundaries of the extremely narrow three-phase alpha-P-delta field are located by the following points: The alpha corner (point 25, fig. 5), as described above; the P corner (point 26, fig. 5) at 8 percent chromium, 34.5 percent nickel, and 57.5 percent molybdenum; and the delta corner (point 27, fig. 5) at 4 percent chromium, 37.5 percent nickel, and 58.5 percent molybdenum.

Some discrepancies were noted in a few alloys of chromium, nickel, and molybdenum near the three-phase alpha-epsilon-sigma field. The microstructure of alloys 475, 569, and 597 showed only two phases, alpha and sigma, although, according to the intended compositions of these alloys, they should contain three phases. Chemical analysis of these alloys moved 569 and 597 into the two-phase alpha-sigma field, thus removing the discrepancy for these two alloys. Alloy 475, however, still remained in the three-phase alpha-epsilon-sigma field. In drawing the phase boundary between the two-phase alpha-sigma and the three-phase alpha-epsilon-sigma fields, alloy 475 was neglected. The microstructure of alloy 640 showed small amounts of the epsilon phase. The chemical analysis of this alloy, however, would place it in the sigma phase field. Both the melted and chemical analyses are given in table V, but the intended melted analysis is plotted in figure 6. Alloys 598 and 600 contained more sigma than would be expected from the phase diagram. This is probably a result of the difficulty in obtaining equilibrium in chromium-rich alloys.

Small amounts of a phase not explainable from the phase diagram were observed in alloys 622 and 447. The identity of this phase is not known. It may have resulted from nonequilibrium or segregation. Small particles of this unknown phase are shown in figure 31, where second-phase particles of alpha in a sigma matrix are also seen.

Alpha boundaries in constant molybdenum sections of chromium-cobaltnickel-molybdenum quaternary system at 1200° C.- Figure 7 is a 1200° C

isothermal diagram in which the alpha phase boundaries for the 2.5-, 5-, 10-, and 20-percent-molybdenum sections through the chromium-cobalt-nickel-molybdenum quaternary system are plotted. Figure 8 is the same diagram with the alloy compositions from table VI indicated.

At constant temperature, a quaternary phase diagram is represented by a tetrahedron, each corner point of which corresponds to 100 percent of one component. The four faces of the tetrahedron, then, are isothermal sections of four ternary systems, and the six edges are isothermal binary sections. In the chromium-cobalt-nickel-molybdenum quaternary system at 1200° C, sections of constant molybdenum content are triangular sections through the tetrahedron parallel to the chromium-nickel-cobalt face. As the amount of molybdenum increases, the size of the equilateral triangle becomes smaller. Because of this change in size, these sections cannot be superimposed upon each other for the purpose of showing, in a two-dimensional diagram, the limits of a phase in the quaternary system. In order to facilitate the diagrammatic presentation of results and to allow direct comparison of the effect of increasing molybdenum content, on the alpha boundary, the following method was used: Each section of constant molybdenum content was projected onto the chromium-cobalt-nickel ternary isothermal section by using the molybdenum corner of the tetrahedron as the center of projection. This projection may be accomplished arithmetically by increasing proportionately the percentages of chromium, cobalt, and nickel in the quaternary composition to add up to 100 percent and then plotting quaternary compositions as if they were ternary chromium-cobaltnickel compositions. The correction factor for each section is the inverted fraction of the total amount of chromium, cobalt, and nickel in the quaternary composition. For example, in a quaternary alloy containing 20 percent molybdenum, 10 percent chromium, 10 percent cobalt, and 60 percent nickel, the chromium, cobalt, and nickel represent eighttenths of the total composition. Multiplying the quaternary percentages of chromium, cobalt, and nickel by the inverted fraction, that is, 10/8, gives the projection of this quaternary point on the chromium-cobaltnickel ternary diagram. While the molybdenum content of alloys used to fix the alpha boundaries was actually never exactly 2.5, 5, 10, or 20 percent, the corresponding boundaries were corrected by factors of 10/9.75, 10/9.5, 10/9, and 10/8, respectively. The results are shown in figure 7, using this method of plotting. An alternative method was used in figure 32 where the actual solubility of chromium in the alpha phase at the various constant molybdenum levels is plotted as a function of actual cobalt content. The data for these curves were interpolated from figure 7 and are tabulated in table XIX. The alpha phase boundary in the 1200° C isothermal section of the chromium-cobalt-nickel ternary system was determined by Manly and Beck (reference 1) and is included for comparison.

As seen from figures 7 and 32, increasing the molybdenum content decreases the solubility of chromium in the alpha phase, thereby decreasing the extent of the alpha field at-1200° C. At the chromium-nickel side of the diagram, the solubility of chromium in alpha first increases slightly and then rapidly decreases beyond about 5 percent molybdenum.

No attempt was made to identify the phases which coexist with the alpha phase in the quaternary system, but some general statements can be made from observations of the chromium-cobalt-molybdenum and chromium-nickel-molybdenum ternary systems at 1200° C. On the chromiumcobalt side of the diagram, the sigma phase coexists with alpha from O to approximately 12.5 percent molybdenum. Between 12.5 and approximately 16 percent molybdenum, the ternary R phase coexists with the alpha phase at the chromium-cobalt side of the diagram. Above 16 percent molybdenum, the mu phase coexists with alpha. On the chromium-nickel side of the diagram, the alpha phase coexists with epsilon from 0 to approximately 3 percent molybdenum and with the ternary sigma phase between 3 and about 19 percent molybdenum. The ternary P phase coexists with alpha on the chromium-nickel side between 19 and 33 percent molybdenum. The delta and quaternary alpha phases coexist over very limited composition ranges, above 33 percent molybdenum. It is not known whether the chromiumcobalt-molybdenum and chromium-nickel-molybdenum sigma phases form an uninterrupted series of solid solutions through the quaternary system at 1200° C, although there is some basis for expecting it. This is treated under Discussion.

In the 2.5-, 5-, and 10-percent-molybdenum alpha boundaries, no sharp breaks were found from the metallographic data, and X-ray parameter plots are not available to check this. However, in figure 32, there is some indication of a cusp on the 2.5 and 5 percent alpha boundaries at approximately 20 percent actual cobalt content. Evidence of two corners on the 20-percent-molybdenum alpha phase boundary was found, one being at approximately 61 percent actual cobalt content (point 28, fig. 32) and the other, at approximately 24 percent actual cobalt content (point 29, fig. 32). The lattice parameters of saturated alpha alloys along this boundary were plotted as a function of actual cobalt content in figure 33. The data for this plot are given in table XX. Two distinct breaks occur in the curve of parameter against composition at approximately 24 and 62 percent actual cobalt content (points 28 and 29, fig. 33). This is in good agreement with the microscopic results.

DISCUSSION

In view of the fact that the face-centered cubic (alpha) solid solutions form the matrix of almost all practically useful high-temperature

alloys, the extent of the alpha phase field in the chromium-cobaltnickel-molybdenum quaternary system and in the four adjoining ternary
systems at 1200° C is of great interest. The alpha phase field in the
1200° C isothermal section of the chromium-cobalt-nickel ternary system
was shown previously to be very broad (reference 1). In the cobaltnickel-molybdenum ternary system, also, the alpha phase field is quite
large. As the cobalt content increases, the solubility of molybdenum
in the alpha phase decreases from 36 percent in the nickel-molybdenum
binary system at 1200° C to 22.5 percent molybdenum in the cobaltmolybdenum system. Both the delta and mu phases, which coexist with
alpha in the cobalt-nickel-molybdenum ternary system at 1200° C, are very
hard, although their presence in alpha alloys does not appear to have
nearly so drastic an effect on toughness as that of the sigma phase.

Solid solutions of the alpha phase surround the cobalt corner of the chromium-cobalt-molybdenum ternary isothermal section and the nickel corner of the chromium-nickel-molybdenum ternary isothermal section. In both of these systems, the extent of the alpha phase field is small in comparison with the chromium-cobalt-nickel and the cobalt-nickel-molybdenum alpha fields. It is interesting to note that in the chromium-nickel-molybdenum isothermal section the sigma phase coexists with alpha above 3.5 percent molybdenum. The fact that the brittle sigma phase may precipitate out of supersaturated alpha containing more than 3.5 percent molybdenum is of great practical importance, since considerable amounts of the sigma phase may seriously impair the forging characteristics of alpha alloys.

In the chromium-cobalt-nickel-molybdenum quaternary system, the extent of the alpha phase field at 1200°C becomes smaller as the molybdenum content increases. In quaternary alloys above 36 percent molybdenum, alpha probably does not occur in a single phase field, although it coexists with other phases up to approximately 61 percent molybdenum.

Some interesting features were observed in the variation of lattice parameter of alpha with composition. Taylor (reference 19) has shown that the general effect of increasing cobalt content in nickel-cobalt binary alloys is to increase the lattice parameter of the alpha phase. Figure 26 shows that the lattice parameter of saturated nickel-molybdenum alpha decreases with increasing cobalt content. Also, from figure 33, along the alpha boundary in the 20-percent constant-molybdenum quaternary section, the lattice parameter again decreases with increasing cobalt content. Molybdenum going into binary solid solution increases the lattice parameter of both nickel and cobalt. Although nickel itself has a smaller ao than cobalt, the lattice parameter of saturated nickel-molybdenum alpha is larger than that of saturated cobalt-molybdenum alpha. Apparently, chromium has the same

general effect as molybdenum. The peculiar variation of the lattice parameter along the alpha boundary in the 1200°C isothermal section of the chromium-cobalt-molybdenum ternary system (fig. 28) may be interpreted as shown in figure 34, which is a schematic sketch of the ternary section with isoparameter lines drawn in the alpha phase field. The decrease of lattice parameter beyond the corner of the three-phase alphasigma-R field results from the intersection of lower and lower isoparameter lines with the alpha phase boundary. At the alpha-R-mu three-phase-field corner, the lattice parameter reaches a minimum value, and beyond this point the alpha boundary again intersects successively higher isoparameter curves until the cobalt-molybdenum side of the diagram is reached.

The accuracy of the location of the alpha phase boundaries is estimated to about 1 percent of any component. Chemical analysis on all definitive boundary alloys gave generally good correlation with intended compositions. Any large discrepancies were usually found to be errors in chemical analysis, although, in some instances referred to in the description of results, the alloy compositions appeared to be in fact different from the intended compositions.

The occurrence of the brittle intermetallic sigma phase in alloy systems involving the transition elements is interesting from both the fundamental and practical viewpoints. The presence of sigma, even in small amounts, in high-temperature alloys is undesirable because of the brittleness which it imparts to alloys. The composition ranges over which sigma exists are, therefore, important. In the cobalt-nickelmolybdenum ternary system at 1200° C the sigma phase does not occur. However, solid solutions of the chromium-cobalt sigma phase penetrate deep into the 1200° C isothermal section of the chromium-cobaltmolybdenum ternary system, and in this system sigma coexists with alpha below 13 percent molybdenum (fig. 3). In the present investigation, a ternary sigma phase was found in the 12000 C isothermal section of the chromium-nickel-molybdenum ternary system. The sigma phase in this system was also found independently by Putman, Grant, and Bloom (reference 20). In a discussion of their paper, Beck (reference 20) presented a tentative diagram for the chromium-nickel-molybdenum ternary system at 1200° C. This diagram was refined in the present investigation and is presented in figure 5. At 1200°C, the sigma phase coexists with alpha in chromium-nickel-molybdenum alloys containing from 3.5 to 19 percent molybdenum.

The sigma phases in the chromium-cobalt-molybdenum and chromium-nickel-molybdenum ternary systems may form a continuous series of solid solutions in the chromium-cobalt-nickel-molybdenum quaternary system at 1200° C. If such quaternary sigma solid solutions do occur, the sigma phase might very likely coexist with alpha in the 1200° C isothermal quaternary section over wide ranges of the alpha phase boundary

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in alloys containing 3.5 to 13 percent molybdenum. Some indication that the sigma phases do form an uninterrupted series of quaternary solid solutions is found in the 10-percent-constant-molybdenum section. The alpha boundary in this section has no pronounced break suggestive of a three-phase-field corner, and sigma does coexist with alpha at both ends of the phase boundary.

The shape and location of the sigma phase field in the chromiumcobalt-molybdenum and chromium-nickel-molybdenum isothermal sections give further support to the previous observation (reference 15) that atoms of transition elements having the same crystal structure can substitute for each other in the formation of the sigma phase. In the chromium-cobalt-molybdenum isothermal section (fig. 3) the sigma phase field extends in a direction parallel to the chromium-molybdenum side of the diagram, and approximately half of the chromium is replaced by molybdenum. Again in the chromium-nickel-molybdenum ternary isothermal section (fig. 5), the ternary sigma phase occurs in a long, narrow field parallel to the chromium-molybdenum side of the diagram. In these two systems, body-centered cubic transition elements are substituting for each other. In the chromium-cobalt-nickel ternary system at 1200°C, the sigma phase field is parallel to the cobalt-nickel side of the diagram, indicating that nickel is replacing cobalt (both face-centered cubic) while the chromium content remains essentially unchanged (references 1 and 15).

As the molybdenum content of the sigma phase increases, the lattice expands in both the chromium-cobalt-molybdenum and the chromium-nickel-molybdenum systems. It is interesting that an X-ray diffraction pattern of a chromium-cobalt-molybdenum sigma alloy saturated with molybdenum is almost identical to a pattern of chromium-nickel-molybdenum sigma of the lowest molybdenum content.

The existence of a ternary sigma phase in the chromium-nickel-molybdenum ternary system is rather unexpected, as none of the adjoining binaries are known to have a sigma phase. From the results of the investigation of the 1200° C isothermal section of the chromium-cobalt-nickel ternary system, Beck and Manly (reference 15) suggested that the nickel-chromium system, which is not reported to include any intermediate phase, might contain a sigma phase at some lower temperature, and Duwez and Baen (reference 16) note that, according to the criteria postulated by them, the nickel-chromium system would be expected to have a sigma phase. The ternary sigma phase found in the chromium-nickel-molybdenum system may have some connection with the suspected nickel-chromium binary sigma phase. Attempts were made to confirm this by annealing for long periods of time an alloy of 38 percent nickel and 62 percent chromium at temperatures below 600° C. These experiments were, however, not successful. This does not rule out the possibility of the existence of sigma

in this system at a temperature so low that the diffusion rates are prohibitively low.

From the experimental results of this and other investigations of alloy systems in which the sigma phase occurs, a criterion for the formation of the sigma phase was developed in terms of electron vacancy concentration in the 3d sub-band. Details of this development are given in reference 21. Briefly, the procedure is to calculate the electron vacancy concentration $N_{\rm V}$ in any alloy from the 3d sub-band electron vacancies in each component of the alloy, and from the atomic percent concentration of the components. Electron vacancy numbers for chromium, cobalt, and nickel were given by Pauling (reference 22; see also reference 23). Molybdenum was assumed to have the same number of vacancies in the 3d sub-band as chromium, on the basis that their electronic structure is the same in the outermost shells. The formula for calculating $N_{\rm V}$ is given as

$$N_v = 4.66(\text{Cr} + \text{Mo}) + 3.66(\text{Mn}) + 2.66(\text{Fe}) + 1.71(\text{Co}) + 0.61(\text{Ni})$$

It was found possible to define fairly well the composition ranges over which the sigma phases occur in terms of electron vacancy concentration calculated from this formula. Almost all of the binary and ternary sigma phase fields were found to fall within a rather narrow range of electron vacancy concentration. The favorable range of concentration seems to be from 3.15 to $3.65 \rm N_V$. It appears that the composition ranges of other intermetallic phases occurring in alloy systems of transition elements can also be defined in terms of electron vacancy concentration, although some overlapping may occur.

There are some interesting features connected with the solid solutions designated mu in the present investigation. For example, the mu phase, commonly referred to as epsilon in the cobalt-molybdenum binary system, is isomorphous with the corresponding intermediate phase in the iron-molybdenum, cobalt-tungsten, and iron-tungsten binary systems (reference 11). Koster and Tonn (reference 24) reported that the mu phases in the iron-molybdenum and cobalt-molybdenum binary systems form a continuous series of solid_solutions across the 1300° C isothermal section of the iron-molybdenum-cobalt ternary system. In the iron-molybdenum-cobalt system, the mu phase field extends in a direction approximately parallel to the iron-cobalt side of the diagram, so that the face-centered cubic elements iron and cobalt substitute for each other while the molybdenum content remains unchanged. As shown in the present work, solid solutions of the mu phase extend more than halfway across the 1200° C isothermal section of the cobalt-nickelmolybdenum ternary system (fig. 1). Here again, the face-centered cubic elements cobalt and nickel replace each other, while the molybdenum

content remains essentially unchanged. The body-centered cubic elements chromium and molybdenum seem to replace each other in forming the mu phase in the 1200°C isothermal section of the chromium-cobalt-molybdenum ternary system (fig. 3), with the cobalt content remaining constant. In agreement with these observations, one would expect that solid solutions of the mu phase in ternary systems of iron, cobalt, molybdenum, and tungsten would also extend in such a manner that atoms of elements having the same crystal structure could substitute for each other. The intermediate nickel-molybdenum delta phase forms only limited solid solutions in the cobalt-nickel-molybdenum and chromium-nickel-molybdenum isothermal sections (figs. 1 and 5).

In addition to the ternary sigma phase, another new ternary phase P was discovered in the 1200° C isothermal section of the chromium-nickel-molybdenum ternary system. The P phase field is located between the ends of the sigma and delta phase fields (fig. 5). In physical characteristics and microstructure, the P and sigma phases are very much alike, but the X-ray diffraction patterns of the two phases are quite different. The differentiation between P and sigma provides a good example of the need to supplement microscopic work with X-ray diffraction to identify the phases. In alloys consisting wholly of P or sigma, it was impossible to identify the phase present by any of the microscopic techniques used in this investigation, even though it was possible to differentiate microscopically between sigma and P when they occurred side by side in the same alloy.

The 1200° C isothermal section of the chromium-cobalt-molybdenum ternary system was also found to contain a ternary phase, located between the ends of the sigma and mu phase fields (fig. 3). This phase R has a distinctive X-ray diffraction pattern (table XIII) and it is easily differentiated microscopically from the sigma or mu phases when these phases occur in the same alloy. However, as in the case of chromium-nickel-molybdenum ternary sigma and P phases, chromium-cobalt-molybdenum alloys consisting wholly of the R or sigma phases could not be differentiated microscopically (except possibly by microhardness). Final identification was made in all cases by X-ray diffraction.

CONCLUSIONS

A survey at 1200° C of portions of the chromium-cobalt-nickel-molybdenum quaternary system and of the component cobalt-nickel-molybdenum, chromium-cobalt-molybdenum, and chromium-nickel-molybdenum ternary systems indicated the following conclusions:

1. A portion of the 1200° C isothermal section of the cobalt-nickel-molybdenum ternary system was investigated, using 47 vacuum-melted alloys with molybdenum contents up to a maximum of 62 percent. The system features an extensive range of face-centered cubic (alpha) solid solutions based on the cobalt-nickel binary system, solid solutions based on the nickel-molybdenum binary delta phase, and solid solutions based on the cobalt-molybdenum binary mu phase. The narrow delta and mu phase fields penetrate into the ternary system to 6.5 percent cobalt and 24 percent cobalt, respectively, and coexist in a narrow two-phase delta-plus-mu field. The alpha phase coexists with the delta and mu phases in two wide two-phase fields and an alpha-delta-mu three-phase field.

- 2. A portion of the 1200° C isothermal section of the chromiumcobalt-molybdenum ternary system was determined, using 85 vacuum-melted alloys with molybdenum contents up to a maximum of 53 percent and cobalt contents of 35 percent and more. The phases present in this composition range were found to be the face-centered cubic (alpha) solid solutions around the cobalt corner, solid solutions based on the chromium-cobalt binary sigms phase, solid solutions based on the cobalt-molybdenum binary mu phase, and a new ternary phase R, of fairly small composition range, which does not appear in any of the binary systems. The sigma phase was found to extend in an elongated field from the chromium-cobalt binary into the ternary system along a line of approximately constant cobalt content of 45 percent, taking molybdenum into solution up to a maximum of 31 percent, so that up to about half of the chromium is replaced by molybdenum. The R phase, comprising a range of chromium contents from about 14.5 to 19.5 percent, is located between the ends of the sigma phase field and of the mu phase field, and it coexists with the sigma and mu phases in 2 two-phase fields. The cobalt-molybdenum binary mu phase extends into the ternary system up to approximately 11 percent chromium. The alpha face-centered cubic solid solutions coexist with the sigma. R, and mu phases in three rather wide two-phase fields and two narrow three-phase fields of alpha, sigma, and R and of alpha, R, and mu.
- 3. The 1200° C isothermal section of the chromium-nickel-molybdenum ternary system was investigated in the composition range of 0 to 59 percent molybdenum and 18 to 100 percent nickel, using 119 vacuum-melted alloys. The following phases were located: The face-centered cubic (alpha) solid solutions around the nickel corner, the body-centered cubic (epsilon) solid solutions based on chromium, solid solutions based on the nickel-molybdenum binary delta phase, the ternary sigma solid solutions which, at 1200° C, are not connected with any of the binary systems, and another ternary phase P, which has not been encountered in any of the binary systems. The ternary sigma phase is located in a long, narrow field between about 21 and 49 percent chromium and approximately 23 to 30 percent molybdenum, which extends parallel to the

chromium-molybdenum side of the ternary diagram, as found also in the chromium-cobalt-molybdenum ternary system. Here, too, the shape and location of the sigma field indicate that chromium and molybdenum can substitute for each other in forming the sigma phase. Successive pairs of the epsilon, sigma, P, and delta phases coexist in 3 two-phase fields. The alpha solid solution coexists with the epsilon, sigma, P, and delta phases in 4 two-phase fields and 3 three-phase fields of alpha, epsilon, and sigma; alpha, sigma, and P; and alpha, P, and delta. The two-phase alpha-plus-sigma field covers an extensive area, starting at molybdenum contents as low as 3.5 percent.

- 4. The alpha solid-solution phase boundary in the chromium-cobaltnickel-molybdenum quaternary system at 1200° C was determined for constant molybdenum contents of 2.5, 5, 10, and 20 percent. The over-all effect of molybdenum additions to the chromium-cobalt-nickel ternary system at 12000 C is to decrease the extent of the face-centered cubic (alpha) solid solutions, that is, to shift the alpha phase boundary toward lower chromium contents. At 1200° C the alpha phase boundary in the 2.5-percent-molybdenum section is essentially the same as that in the chromium-cobalt-nickel ternary system, except near the chromiumnickel side, where it is moved toward higher chromium contents. phases which coexist with the alpha phase in the 2.5-percent-molybdenum section are sigma and epsilon. In the 5-percent constant-molybdenum section, the alpha-phase-boundary shift becomes more evident, although the magnitude of the shifting is still relatively small. The sigma phase is seen to coexist with the alpha solid solutions near both the chromium-cobalt and the chromium-nickel side of the diagram. due to the presence of the ternary sigma phase in the chromium-nickelmolybdenum ternary system at 1200° C. The alpha phase also coexists with epsilon and sigma in a three-phase field near the chromium-nickel side of the diagram. The shift of the alpha phase boundary toward lower chromium contents is more pronounced in the 10-percent-molybdenum section of the isothermal quaternary system. Again the sigma phase is seen to coexist with the alpha solid solutions at both the chromium-cobalt and the chromium-nickel ends of the alpha phase boundary. The alpha phase also coexists with epsilon and sigma in a three-phase field near the chromium-nickel side of the diagram. The alpha phase boundary in the 20-percent-molybdenum section is shifted to 4 percent chromium on the chromium-cobalt side of the diagram and to 30 percent chromium on the chromium-nickel side. The mu phase coexists with the alpha solid solutions near the chromium-cobalt side of the quaternary section, whereas the ternary P phase coexists with the alpha phase at the chromium-nickel end of the alpha phase boundary.
- 5. The results of this investigation give further support to the conclusion that atoms of both face-centered and body-centered cubic transition elements are required to form the sigma phase and that

elements having the same crystal structure can substitute for each other. It appears that the same criterion may hold for ternary solid solutions of the cobalt-molybdenum intermediate mu phase, which is isomorphous with the corresponding intermediate phases in the iron-molybdenum, cobalt-tungsten, and iron-tungsten binary systems. The composition ranges over which the sigma phases exist can be described fairly well in terms of electron vacancy concentration in the 3d sub-band. This suggests a criterion for predicting whether or not the sigma phase may occur within any given composition range in alloy systems involving the transition elements.

University of Notre Dame
Notre Dame, Ind., January 31, 1951

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TABLE I

LOT ANALYSES BY WEIGHT PERCENT

Element or compound	Cobalt rondelles	Chromium	Nickel	Molybdenum
C CaO Co Cr Cu Fe Ho Mn Mo No No Pb SiOo	0.17 .12 Bal. .02 .12 .04 .06 .46 	0.01 Bal. Trace .06 .045 .010 .51 .001 .012	0.6 to 0.8 .01 .01 .01 .01 .01	0.003 max005 max. Bal003 approx.

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TABLE II
CRUCIBLES USED FOR MELTING

Zirconia	Stabilized zirconia	Alundum
273	323	272
311 to 314	330 to 332	280
320	354	288
322	360 to 362	289
402 to 405	366	296 to 298
414 to 419	376 to 383	321
		370 to 372
		375
		384 to 401
		407 to 413
		420 to 715

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TABLE III DATA FOR THE PHASE DIAGRAM OF THE COBALT-NICKEL-MOLYBDENUM SYSTEM AT 1200° C

Alloy	(we	Analysis		Phase	Phases found from - (percent) (percent) Microstructure						Annealing time
	Co	Ni	Мо	Alpha	Delta	Mu	Alpha	Delta	Mu	X-ray	(hr)
	<u> </u>			<u>L.</u>	Alpha	boundary	alloys ^a	L	1	1	
haaa		l	00.00		T			Γ		1	10
ъ399 400	77.08	65	22.92 35	98 100		2	95 100		5	Alpha Alpha	. 48 48
401	30	40	36	98		2	100		Trace	Alpha	48
b 407	20	50	30	100			100			Alpha	56
	17.4	50	32.6	i .	l	ĺ	1		Ì	1	-
b408	49.62	24.95	25.43	99		1	100		Trace	Alpha	65
426	32	¥2 68	26	100			100			Alpha	48
427 432	79		32 21	100			100 100			Alpha Alpha	48 148
440	50	26	22	100			100			Alpha	₩A
441	20	57	23	100			100			Alpha	48
D461	49.89	26.47	23.64	100			100			Alpha	48
462	20	54	26	100			100			Alpha	55 45 45 45 45 45 45 45 45 45 45 45 45 4
ъ486 -		62.92	37.08	97		3	95	5		Alpha	65
1:98	20	53 12	27	100			100			Alpha	48
540	65	12	23	100			100			Alpha	48
541	10 64	58 11	32 25	100 98		2	100 96		4	Alpha	48 L0
552 553	10	5 6	34	100			100			Alpha Alpha	40 ke
553 620	30.24	40.84	28.92	100			100			Alpha	48
627	10.2	53.72	36.08	98	Trace	2	96	1	lз	Alpha	48
679	15	52	33	100		Trace	100		Trace	Alpha	48
		<u> </u>	<u> </u>	<u> </u>	Alph	a-plus-mu	alloys ^c	ł	!	1 6	
421	55		45	30		70	25		75	Alpha plus mu	95
442	55 15 25	35	50	40		60	න 50 න න න		50	Alpha plus mu	95
443	25	35 25	50	30		70	25	l	75	Alpha plus mu	95
444	35 18.28	15	50	20		80	25		75	Alpha plus mu	95
^b 536	18.28	24.12	57.60 40	1	{	99	5		50 75 75 95 20	Mu	95
546	14	46	40	75		25	80			Alpha plus mu	95
547 548	18 18	42 38	40 40	70		30	12		25	Alpha plus mu	95
554	22	26 28	50	70 35		30 65	١١٤		25 95	Alpha plus mu Mu	92 05
715	12	48	40	35 80		ဆိ	75 75 5 75		25	Alpha plus mu	95 95 95 95 95 95 95 95
			l			Mu alloy	sc				
D550	22.74	18.77	58,49			100			100	Mu .	95
596	23	16	61			100			100	No.	95
596 629	20	20	60			100			100	Mu	95 95 95
			l		Alpha-,	olus-delt	s alloys ^c			<u> </u>	
	-	49	51	40	60		50	50		Alpha plus delta	Q5
hih					. ~						22
414 465	5	145	50	l hs	55		śŏ	50	l	lalpha plus deltai	95
465	5 k	45 38	50 58	45 10	55 90		50	50 50 85		Alpha plus delta Delta	95 95
465 520	. 4	45 38	50 58 58	45 10 10	55 90 90	Trace	50 15 12	85 85		Alpha plus delta Delta Delta	95 95 95
465	5 4 7	45 38 35 36	50 58 58 60	10	90	Trace	50 15	50 85 85 100	3	Delta	95 95 95 95 95
465 520 521	ች 7	45 38 35	50 58 58	10 10	90 90 99	Trace	50 15 12 Trace	85 85	3	Delta Delta	95 95 95 95
465 520 521 677	7.05	45 38 35 36 30.78	50 58 58 60 62.17	10 10	90 90 99 Delta	 a-plus-mu	50 15 12 Trace	85 85 100		Delta Delta Delta Delta	
465 520 521 677	ች 7 ት	45 38 35 36	50 58 58 60	10 10 1	90 90 99 Delta 95 98	 a-plus-mu 5 2	50 15 12 Trace alloys ^c	85 85 100 95 95	3	Delta Delta Delta	95 95 95 95 95
9595 9628	7.05	45 38 35 36 30.78 32.65	50 58 58 60 62.17	10 10 1	90 90 99 Delta 95 98	 a-plus-mu 5 2	50 15 12 Trace alloys ^c 1	85 85 100 95 95	5 4	Delta Delta Delta Delta Delta	
465 520 521	7.05	45 38 35 36 30.78	50 58 58 60 62.17	10 10 1	90 90 99 Delta 95 98	 a-plus-mu 5 2	50 15 12 Trace alloys ^c	85 85 100 95 95		Delta Delta Delta Delta Alpha plus delta	95 95 95
9595 9628	7.05	45 38 35 36 30.78 32.65	50 58 58 60 62.17 60.13	10 10 1	90 90 99 Delta 95 98	5 2 -delta-pl	50 15 12 Trace alloys ^c 1	85 85 100 95 95	5 4	Delta Delta Delta Delta Delta Delta Delta Delta Delta	95 95 95
465 520 521 677 595 628 464 522	7.05 7.22	45 38 35 36 30.78 32.65	50 58 58 60 62.17 60.13	10 10 1 1	90 90 99 Delta 95 98 lphs-plus	5 2 -delta-pl	50 15 12 Trace alloys ^c 1 us-mu allo	85 85 100 95 95 95	5 4	Delta	95 95 95
165 520 521 677 595 628	7.05 7.22	45 38 35 36 30.78 32.65	50 58 58 60 62.17 60.13	10 10 1 1	90 90 99 Delta 95 98 · 1phs-plus 35 60	 a-plus-mu 5 2 -delts-pl	50 15 12 Trace alloys ^c 1 us-mu sllo	85 85 100 95 95 95 95 30 70	30	Delta Delta Delta Delta Delta Delta Delta Delta Delta	95 95

All forged prior to annealing.

Chemical composition from analysis. Composition given for other alloys is the intended melted analysis.

Chot forged prior to annealing.

TABLE IV DATA FOR THE PHASE DIAGRAM OF THE CHROMIUM-COBALT-MOLYBDENUM SYSTEM AT 1200° C

Alloy		nalysis		Pha	ses fro		e 4				ercent)		Annealing time
	Cr	Co	Мо	Alpha	8igma	R	Mu	Alpha	Sigms	ructure R	Mu	X-ray	(hr)
 	- CF		MO	итрпа	O18mm		Mu	ктриа	DIE				L
	,	. —				Alp	ha boun	dary al	loys ^a				·
272 366	33.15 29.45	64.35 65.55	2.5	90 100	10			85 100	15			Alpha Alpha	48 48
b378	30.31	65.96	3.73	100				100				Alpha	表记证方表表表表表表表表表表表表表表表表
b399	l	77.08	22,92	98			2	95			5	Alpha	¥8
413	10	72	18	95		j	5	90			10	Alpha	148
430 431	10 20	76 70	14 10	100				100				Alpha Alpha	40 10
432		79	21	100				100				Alpha	48
453	20	79 68	15	100		l		100				Alpha	48
455	10	74	16	100]]	100				Alpha	48
456	3	77	20	100				100				Alpha	1 48
^b 471 487	28.83	64.8	6.37	98 100	2	{ 	!	98 100	2			Alpha	48
b499	25.61	76 65.4	8.99	100			Trace	100			Trace	Alpha Alpha	16
606	22.5	64.5	13	90	10			90	10			Alpha	148
9619	14.58	69.78	15.64	100				100				Alpha	48
b632	9.88	72.99	17.13	100			[']	100				Alpha	48
644	25	64.5	10.5	92	8			95	5			Alpha	48
b645	17.42	68.01	14.57	100				100				Alpha	148
b665	14.7	68.65	16.65	95			5	98			2	Alpha	40
				_		Alph	e-plus-	sigma a	lloys ^c		<u></u>		
273	31.28	60.72	8	75	25			65	35			Alphe	96
268	38.02	59.48	2.5	75 65	35			60	40			Alpha plus sigma	96
314	36.1	58.9	5	62	35 38			60	140			Alpha plus sigma	96
322	36.8	55.2		50	50			50	50			Alpha plus sigma	105
b330	34.16	63.41	2.43	87	13			90 40	10 60			Alpha	56 96
384 1398	42.3 31.7	51.7 63.09	5.21	35 87	65 13			90	10			Sigma plus alpha Alpha	96 96
470	29	62	9.21	85	15			80	20			Alpha	145
0474	32.22	47.57	20.21	15	15 85 55 60			15	85			81gma	245
5i2	27	53	20	45	55			50	50			Alpha plus sigma	96
538	29.5	51.5	20	40	60			50	50			Sigma plus alpha	96 96
539	38	48	14	20	80			25	75			Sigma	96
543 555	28 41	46 45	26 14	5 5	95			5 10	95 90			Signa Signa	145 150
9557	23.3	56.08	20.62	55	95 45			50	50	nace		Sigma Alpha plus sigma	96
b616	33.75	44.98	21.27	75	95			75	95			Signs	150
Þ682	7.2	45.6	47.2	5	92		J	10	90			Signa	150
	1		l			A1;	pha-plus	Ralle	oys ^c				
D412	19.36	65.29	15.35	90		10		90		10		Alpha	76
533	19.30	52	32	40		60		50		10 50		Alpha plus R	96
b558	19.81	57.9	22.29	55		45		50		50		Alpha plus R	96
560	16	60	24	60		40		60		40		Alpha plus R	96
592	18	ի կ	38	2		98]	2		98		R	150
						Alp	na-plus-	-mu all	OAR _C			!	
b534	11.6	54.27	34.13	45			55	50			50	Alpha plus mu	96
535	8	56	36	50			50	50			50	Alpha plus mu	96
535 5544	7.86	43.05	49.09	50 5 65			95	5			95	Mu	96
Þ573	12.7	61.31	25.99	65)		35	75			25	Alpha plus mu	96

SAll forged prior to annealing.

bChemical composition from analysis. Composition given for other alloys is the intended melted analysis.

cNot forged prior to annealing.

TABLE IV - Concluded

DATA FOR THE PHASE DIAGRAM OF THE CHROMIUM-COBALT-MOLYBDENUM SYSTEM AT 1200° C - Concluded

		nalysis ht perc		Pha	ses from		e 4		Phe	ses fou (perc	nd from		Annealing
Alloy			,							ructure		X-ray	time (hr)
	Cr	Co	Мо	Alpha	Sigma	R	Mu	Algha	Signa		Mu	<u> </u>	<u> </u>
<u> </u>	,				Alp	ha-plus	-sigma-	plus-R	alloys			,	,
5513 556 605 633 649	19.88 24 18.5 22.5 24.5	49.78, 50 45.5 46.5 47	30.34 26 36 31 28.5	20 18 8 5 5	5 80 2 70 90	75 2 90 25 5		20 20 9 10 5	10 80 1 70 90	70 Trace 90 20 5		R plus alpha Sigma R Sigma plus R Sigma	96 96 96 96
	<u> </u>	<u> </u>	<u> </u>	<u>. </u>	A1	pha-plu	s-R-plu	s-mu el	loys ^c		L		
b559	13.67	53 12	33.21	140		·	1	40		60	Trace	R plus Alpha	96
585 680 681	13.76	53.12 60.99 44 43.03	25.25 44.5 47.15	70 2 4		59 10 43 Trace	20 55 96	75 2 2		10 48 Trace	15 50 98	Alpha plus mu Mu plus R Mu	
	1,,,,,	1.504,5	1 .1	<u> </u>	<u> </u>	<u> </u>	igma al	loysc					
392	52.6	41.4	6		100			Ī	100			Sigma	150
393 423 b472 473 b485 b500 530 b531 543	56.4 34 39.97 37 33.99 41.3 32 28.37 28	37.6 36.56 43.56 42.94 38.56 40.69 46	6 30 19.47 20 23.07 20.14 26 30.94 26		100				80 75 100 100 100 95 100 495 95			Sigma Sigma Sigma Sigma Sigma Sigma Sigma Sigma	150 150 150 150 150 150 150 150
590 593 594 624 625	46 27.1 30 28.5 27	42.4 42.4 42.4 44.5 45.5	10 30.5 28 27 27.5	=======================================	100 100 100 100 100			Trace	100 100 100 100 100	Trace		Sigma Sigma Sigma Sigma Sigma	150 150 150 150 150
						S1gma	-plus-R	alloys	c				
6422 6497 532 650	24.42 19.79 24 19	45.82 41.95 42 44	29.76 38.26 34 37		85 10 60 15	15 90 40 85			80 10 50 25	20 90 50 75		Sigma R Sigma plus R R	150 96 96 96
							R alloy	₈ C					
b514 b591	15.71 13.78	41.69 43.89	42.60 42.33			100 100				100 100		R R	96 96
						R-p.	lus-mu	alloysc					
515 666	12 16	42 40	47 44			40 98	60 2			50 95	50 5	Mu plus R R	96 96
							Mu allo	ysc					
516 545 5574 5630 631	6 11 8.51 9.4 10	41 38 40.37 42.8 42	53 51 51,12 47,8 48	2		2	100 99 100 98 98	2		2	100 96 100 98 98	Mu Mu Mu Mu Mu	96 96 96 96 96

Chemical composition from analysis. Composition given for other alloys is the intended melted analysis.

CNot forged prior to annealing.

dalso two second phases.

₹, ₫

		nalysis ht perc			Phases (r	from fi						(perce	d from	-	Annealing
Alloy	Cr	N1	Мо	Alpha	Epsilon	Sigme	P	Delta	Alpha	Micro Epsilon	Signa	P	Delta	X-rey	time (hr)
			1		1-	<u> </u>	L	Alpha	alloys	a		·	<u> </u>	!	<u> </u>
200		0								T		[1-	43-1-	48
b360	42.51 39.9	55.08	2.41	100					100					Alpha Alpha	48
375 1382	40.56	55.1 54.92	4.52	100					100					Alpha	48
400		65 68	35	100					100					Alpha	48
427		68 64	32 26	100					100					Alpha Alpha	48 48 48 48 48
433 434	10 26	58	16	100					100					Alpha	48
P437	35.99	54.28	9,73	100					100		Trace			Alpha	48
P438 .	25.48	157.31	17.21	100		Trace			100		1			Alpha	148
454 10486	10	62.5 62.92	27.5 37.08	100					100					Alpha Alpha	48
₽688	5	61.75	33.25	100				Trace	100				Trace	Alpha	48
							Alph	a-plus-	epsilon	alloysc				•	<u> </u>
296	56.33	41.17	2.5	25	hs.	T	T		50	50				Alpha plus epsilon	96
9311	43.37	54.17	2.5	22 25 26 25 25 25 25 25 25 25 25 25 25 25 25 25	45 5				95	5				Alpha	96 48
9387	53.86	40.28	5.86	50	i 5α				50	50				Alpha plus epsilon	96 96 96 96 96
404	70.5	23.6	6	2	95 95 65 70				10 10	90 90				Epsilon Epsilon	96
449 562	60 60	30 35	10	35	¥2 65				40	60				Epsilon plus alpha	96
563	57.5	35 35	5 7.5	36	70				30	70				Epsilon plus alpha	96
614	57.5 43.5	53	3.5	98	2.				95	5				Alpha	48
							Alph	a-plus-	sigme a	lloysc			-		
389	42.75	52.25	5	95		5			95		5			Alpha	48
410	26	54	20	95 85 85		15	~		90 80		10			Alpha	26
411 1-10	40	50	10 20	85		15			80		20			Alpha Sigma plus alpha	20
418 6435	45 36.25	35 53.63	10.12	8		75			95		75 5	~		Alpha	56 56 96 48
445	28	149	23	75		25 65			75		25			Alpha plus sigma	96
446	32	38 30 46	130	25 98 75 35 5		65			8 8 8 8 8 8 8 8		2555 1559 1959 1959			Signa plus alpha	96
447	40	30	30 18	76		95 30 35 55 60 80			2		92			Sigma Alpha plus sigma	96
451 452	36 43 40	(4 4	13	65		35			60		40			Alpha plus sigma	96
476	40	40	20	45		55			50		50			Sigma plus alpha	96
477	35 48	40 32	25	40 20		80			50 25		50 50 75 80			Sigma plus alpha Sigma	96
478 479	40 41	33	20 26	20		80			20		80	~		8igma	96
480 I	36	33 34	30 40	20		80			25		75			8igmas	96
482	26	32	40	10		90 90			10 5		90			Sigma	96
483 484	33 27	32 38	35 35 25 16.5	10 30		70			140		95 60			Signa Signa plus alpha	96
491), E	30	25	10		90			15		85			Sigma	96
⁵ 569 570	45.98 46.5 44.35	37.87	16.5	40		60			50 40		50 60			Sigma plus alpha	96
570 9507	46.5 հե շա	36 44.67	17.5	30 65		70			60		90 40			Sigma plus alpha Alpha plus sigma	96
608 608	23.5	39	37.5	36		35 70			25		75			Signa plus alpha	96
615	23.5 48.5	30.5	21	30 10		90			25 15		85			61gma	96
617	24.5	32 48	43.5 28.7	65		93			5 70		95 30			Sigma Alpha plus sigma	96 96 96 96 96 96 96 96 96 96 96 96 96 9
634 6641	23.3 22.76	31.6	45.64	5		95		,	5		95			Signa Signa	96
	35	30 48.5	35	1		99			Trace		95 100			Sigma.	150
654 663	22.5	48.5	89 46	75 5		99 93 95 95 95 95 95 95 95 95 95 95 95 95 95			75 5		25 95			Alpha plus sigma	150 96 96
684 687	22 29.5	32 30.5	46	2		%			2		99			Sigma Sigma	96
	۷.,	30.0						pha-plu	· -	ovec.					
409	10	60	30	95					93			7		Alpha	48
415	20	40	40	35			5 65		20			70		P plus alpha	96 48 48
457	22.9	56.45	20.65	90			10		95			5 2		Alpha	48
1460 466	14.28 20	59	26.72	35 90 95 75 80			5 25		95 98 80			20		Alpha Alpha pius P	96
467	10	50 55 45	35 45	ao			20		75 50			25		Alpha plus P	96 96 96
468	10	45	45	35			65		50			25 50 5		P plus alpha	96
b469	22.09	57.33	20.58	95			5		95 3			97		Alpha P	45 06
510 579	10	35 33.72	55 51.21	3			5 95 97 97		5			95		P	96 96
612	15.07 8.5	35 (-	56.5	1			97		5			95		P	96 48
P636	11.26	35 59.46 48.23	29.26	93 65			7		95 70			95 95 5 30		Alpha	48 04
669 689	21.25 6.41	48.23 53.48	30.52 40.11	65 65			35 35		70 70			30 30		Alpha plus P Alpha plus P	96 96
909	0.41	25.40	40.11	62			39	! i	10			. تح		vrhue brue	70

*All forged prior to annealing. Differential composition from analysis. Composition given for other alloys is the intended melted analysis. CNot forged prior to annealing.

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TABLE V - Concluded DATA FOR THE PHASE DIAGRAM OF THE CHRONIUM-NICKEL-MOLYEDENUM SYSTEM AT 1200° C - Concluded

	(wei	Analysi ght per	s cent)	_	Phases f	rom fig	are 6						found percent	from -	Annewling time
VITOA		,								Micro	structu	re		X-ray	(hr-)
	Cr	Mi	Mo	Alpha	Epsilon	Signa	P	Delta	Alpha	Epsilon	Sigma	P	Delta		
								Alpha	-plus-de	lte allo	ysc .				-
414		49	51	40				60	50				50	Alpha plus delta	95
691 692	4.5 4.5	49 54.5 46	51 41 49.5	70				30 60	50 75 50				50 25 50	Alpha plus delta	95 96 96
092	4.5	140	49.5	_~			لتتت						_ >0	Delta plus alpha	70
				,—			Alph	1-plus-		plus-sig		yec .			
320 5448	43.7 48.66 49.82 47 54.5 52.5 55 51 52.6 55.85 48.5	31.25	20.09	90	Trace	10 80			90	1 10	80			Alpha Signa	75 150 150 150 150 150 150 150 150 150 15
150	49.82	37.19	12.99	15 35 35 35 36 36 36 36 36 36 36 36 36 36 36 36 36	36	l 4o			150000000000000000000000000000000000000	25	¥0 50			Signa plus epailon	150
2450 475 564 565 586 588 600 622 655	147	35	13 10.5	1 73	50	\$5.50 \$5.00			20	55	20			Signa plus signa Epsilon plus signa	150
565	52.5	134,54	12.96	ão.	50 40	₩ŏ			36	55 35 50 20	20 35 35 35 75 75 20 20 20 20 20 20 20 20 20 20 20 20 20			Epsilon plus signa	150
566 586	55	30 29 28.73	15 20	10	50 20 38 75 34 10	¥0			20	50 20	30			Epsilon plus signs Signs	150
÷886	52.6	a8.73	18.67	1 2	36	66			5	20	75			8igme	150
7598	55.85	35.1 45.4	9.05	23 65	75	2			30	50 30	20			Epsilon plus sipha plus signa	150
622	50.5	(28)	22	Trace	10	90			50	10	20 85 5			Alpha plus epsilon plus signa Signa	150
655	50 43.5 59.5	52.25	4.25	94	1	5			95	Trace 80	20			Alpha	75
672	29.5	29	11.5	2	93	5			Trace		L			Epsilon plus signa	150
								tope pl	us-sign	-plus-P				·	
568	20	35 32.5	45 48.2	30		30 5	40		30		30 15 95 10	40		P plus signs	96 96 96 96
613 642	19.3 21.25	32.5	46.75	30 5 5		95	90 Trace		10		35	75 Trace		P Sigma	96
642 669	22	32 49	29	76		95 15	15		15		íó	15		Alpha	96
			<u> </u>				All	pha-plu	s-P-plu	-delta a	lloysc	-	٠	<u>' — — — — — — — — — — — — — — — — — — —</u>	
635 670	7	45	48	38			60	2	5 0 8			50 87	Trace	P plus slphs	96 96
670	7.1 5.5	35.8 54	57.1 40.5	38 5 70			70 10	25	70			10	20) P	96 96
690	2.5_) -	+0.5					1						Alpha plus delta	30
								<u> </u>	psilon						
397 599 618	76.2 65 61	17.8	6		100	[-				100	{	[·		Epsilon	150 150 150
618	8	22 27	13 12		98	2				100	Trace			Epsilon Epsilon	150
676	65	25.85	9.25		100				Trace	100				Epsilon	150
								Bpsilo	n-plus-	sigma all	Oysc				
119 567 587 610	61 52.5 51.99 47.6	19 25 27.19 27.2	20 22.5							65 30 30	35 70 70			Signa plus epsilon	150 150 150
201 2587	51.99	27.19	20.82		35 25 2	65 75 98				30 30	70			Signs Signs	150
640	47.6	27.2	25.2 23.5		2	98		[5	95			Signs.	150
6¥3	199 59	27.5	23.5		80	20									1 ~~
	1	1								60	40			Epsilon plus signs	
										60	40	==		Epsilon plus signa	150
1.00		1-4			·				Sigma a						150
416 481	34 37	26 28	40 35			100			Bigma a		100			Signe	150
416 481 493	34 37 47	26 28 23	40 35 30	===		100			Sigma a		100	{		Signa Signa Signa	150
126 181 193 508 509	34 37 47 32 25,06	123	40 35 30 45 48 82		===	100			Bigma a		100			Sigma Sigma Sigma Sigma	150
416 481 493 508 509 511	34 37 47 32 25.06 38	23 26.12 26.12	40 35 30 45 48.82			100			Sigma a	lloysc	100 100 690 690 695 695			Signa Signa Signa Signa Signa Signa	150
481 481 508 509 511 527	34 37 47 32 25.06 38	23 26.12 26.12	40 35 30 45 48.82 40			100 100 96			Sigma a	lloysc	100 100 490 490 490 490 100			Signa Signa Signa Signa Signa Signa Signa	150
416 481 493 508 509 517 578	34 37 47 32 25.06 38 31 31.5 30.5	23 26.12 26.12 24.5 24.5	\$0 35 30 \$5 \$8.82 \$0 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4			100 100 98 100 100			Sigma a	lloysc	100 100 490 490 490 295 490 100 100			Sigma Sigma Sigma Sigma Sigma Sigma Sigma Sigma Sigma	150
161 181 181 181 181 181 181 181 181 181	34 37 47 32 25.06 38 31.5 30.5 25	23 26.12 26.12 24.5 24.5	\$0 35 30 \$5 \$8 \$0 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4 \$4			100 100 98 100 100 100			Biguna a	lloysc	100 100 490 490 490 100 100 100			Signa Signa Signa Signa Signa Signa Signa Signa Signa	150
416 481 493 508 509 517 578 603 604	32 25.06 38 31.5 30.5 25 36 45	23 26.12 26.12 29 24.5 26.5 27.5	35 30 45 82 40 44 43 47 57			100 100 96 100 100 100 100 100			Bigma a	lloysc	100 100 490 490 490 100 100 100 100			Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa	150
\$16 \$81 \$93 508 509 511 527 577 602 603 604 685	34 37 47 32 25.06 38 31 31.5 30.5 25 36 45 22.5	23 26.12 26.12 24.5 24.5	40 35 30 48 40 44 47 5 40 30 47 5			100 100 98 100 100 100 100				lloysc	100 100 490 490 490 100 100 100 100 100 100			Signa Signa Signa Signa Signa Signa Signa Signa Signa	
685	32 25.06 38 31.5 30.5 25 36 45 22.5	23 26.12 26.12 29 24.5 26.5 27.5 24 25 30	35 30 45 88 82 40 44 43 47 5 30 47 5			100 100 98 100 100 100 100 100				lloysc	100 100 490 995 490 100 100 100 100			Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa	150 96 96 96 96 96 96 96 96 96 96 96 96 96
528	32 25.06 38 31.5 30.5 25 36 45 22.5	23 26.12 29 24.5 26.5 27.5 24 25 30	35 30 45 88 82 40 44 43 47 5 30 47 5			100 100 98 100 100 100 100 100				lloysc	100 100 490 490 100 100 100 100 100	50		Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa	150 96 96 96 96 96 96 96 96 96 96 96 96
685	32 25.06 38 31.5 30.5 25 36 45 22.5	23 26.12 26.12 29 24.5 26.5 27.5 24 25 30	35 30 45 82 40 44 43 47 57			100 100 96 100 100 100 100 100	50 20 90			lloysc	100 100 490 995 490 100 100 100 100			Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa	150
528 686	32 25.06 33 31.5 30.5 25 36 45 22.5	23 25.12 22.22 24.5 26.5 27.5 24 25 30	35 30 45 88 82 40 44 43 47 5 30 47 5			100 100 100 100 100 100 100 100 100	5°C 2°C		on -plus	lloysc	100 100 490 490 100 100 100 100 100 100	50		Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa	96 96 96 96 96 96 96 96 96 96 96 96
528 686	32 25.06 33 31.5 30.5 25 36 45 22.5	23 25.12 22.22 24.5 26.5 27.5 24 25 30	35 30 45 88 82 40 44 43 47 5 30 47 5			100 100 100 100 100 100 100 100 100	5°C 2°C		on -plus	lloys ^c	100 100 490 490 100 100 100 100 100 100	50		Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa	96 96 96 96 96 96 96 96 96 96 96 96
528 686 492	32 25.06 38 31 31.5 30.5 25 45 22.5	23 26.12 26.12 22 28 24.5 26.5 27.5 24 25 30	35 30 48.82 40 44 43 47.5 30 47.5			100 100 100 100 100 100 100 100 100	50 20 90		m-plus P el	lloys ^c	100 100 490 490 100 100 100 100 100 100	50 96		Signa	96 96 96 96 96 96 96 96 96 96 96 96 96 9
528 686 492 580	32 25.06 38 31 31.5 30.5 25 36 22.5 22 23.5 20	23 26.12 22 29 24.5 26.5 27.5 24 25 30 28 27.5 30	35 30 45 48.82 40 44 43 47.5 40 30 47.5 50			100 100 100 100 100 100 100 100 100	50 20 90	S1g	m-plus P el	P alloys	100 100 490 490 100 100 100 100 100 100	50 96	60	Sigma Figma	96 96 96 96 96 96 96 96 96 96 96 96 96 9
528 686 492	32 25.06 38 31 31.5 30.5 25 45 22.5	23 26.12 26.12 22 28 24.5 26.5 27.5 24 25 30	35 30 48.82 40 44 43 47.5 30 47.5			100 100 100 100 100 100 100 100 100	50 20 90		m-plus P el	P alloys	100 100 490 490 100 100 100 100 100 100	50 96	 	Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa Signa P	96 96 96 96 96 96 96 96 96 96 96 96 96 9

bChemical composition from analysis. Composition given for other alloys is the intended melted analysis. Got forged prior to annesling.

Also 10 percent of a second phase.

Salso two second phases.

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TABLE VI DATA FOR THE PHASE DIAGRAM OF THE CHROMIUM-COBALT-NICKEL-MOLYBDENUM SYSTEM AT 1200° C

	(Tare	Anal	ysis percen	+1	}	hases :	e 8			found i		Annealing
Alloy	, "`	10110	per our	٠,		(perce	n t)	Mic	rostru	cture		time (hr)
	Cr	Co	Ni	Мо	Alpha	Sigma	Epsilon	Alpha	Sigma	Epsilon	X-ray	(1117)
				2.5	perce	nt-mol	ybdenum (quateri	nary a	lloysa		
280	42.9	16.58	38.02	2.5				c99				48
288	38.02	59.48		2.5	65	35		60	40	ļ	Alpha plus sigma	96
289	35.1	62.4		2.5	85	15		90	10		Alpha	96 48
296	38.02	44.85	14.63					c95				48
297	44.35	15.6	37.05					°95				48
p311	43.37		54.17	2.46	95		5	95		5	Alpha	48
312		31.2	25.35	2.5		ļ - - ;		^{C85}				48
313	44.85	9.75	42.9	2.5				°90)		48
323	39	31.2	27.3	2.5				°98				48
^b 330		63.41		2.43	87	13		90	10		Alpha	48
	36.84		15.61					c99		!		48
332	43.38	9.75	43.87					^c 98				48
b354	37.34	31.90	28.08		100			100			Alpha	48
b360	42.51		55.08	2,41	100			100		1	Alpha	48
b361	41.94	9.51	46.12		100	<i>-</i>		100			Alpha	48
362		24.38		2.5	100	[- -		100			Alpha	48
372		24.38	32.66	2.5				C98		¹. 		48
376	38.76		27.54					^C 99		·		48 48
377	43.39	9.75	44.36	لستيا				c99				40
[5-pe	rcent-	molybo	lenum qua	ternar	y allo	ува		
314	36.1	58.9		5	62	38		60	40	- -	Alpha plus sigma	96
320	43.7		51.3	5	90	10	Trace	90	9	1	Alpha	75
321		19	32.3	5			11000	c90			Arpna	75 48
		65.55		5	100			100			Alpha	48
375	39.9		55.1	5	100			100			Alpha	48
Þ378		65.96		3.73	100			100			Alpha	48
D379		42.49	18.72	4.17	100			100			Alpha	48
15380 l	40.37		37.84					c95				48
b381	39.69	9.34						c98				48
b382	40.56		54.92		100			100			Alpha	48
Þ388	38.84	10	46.56		100			100			Alpha	48
389	42.75			5	95	5		95	5		Alpha	48
	39.06	17.14			100			\mathbf{d}_{100}				48
8esa		63.09			87	13		90	10		Alpha	96
b428			17.16	5.22				°98				48
b429		27.83	29.62					c95				48
436	37.05	28.5		5	100			100			Alpha	48

^aAll forged prior to annealing.

^bChemical composition from analysis. Composition given for other alloys is the intended melted analysis.

^cRemaining percentage is unidentified second phase.

^dAlso trace of unidentified second phase.

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TABLE VI - Concluded

DATA FOR THE PHASE DIAGRAM OF THE CHROMIUM-COBALT-NICKEL-MOLYBDENUM SYSTEM AT 1200° C - Concluded

	(10	Anal;		+)		ases f		Pha		ound from	n -	Annealing
Alloy		CIBILG	Per cen	.0)		(perce	nt)	Mic	crostr	ucture	<u> </u>	time (hr)
	Cr	Co	Ni	Мо	Alpha	Sigma	Epsilon	Alpha	Sigma	Epsilon	X-ray	, ,
			:	10-perc	cent-m	olybdeı	num quate	rnery	alloys	₃ a		
431 b435 b437 506 507 b517 b518 b519 b621 b623 b653	20 36.25 35.99 28.8 34.2 26.75 28.72 31.5 29.48 27.84 33.4	18 33.90	26.87 40.5		100 98 100 100 100 100	2		100 98 100 c90 c85 100 100 100 c95 c93 c92	5	1-	Alpha Alpha Alpha Alpha Alpha Alpha 	48 48 48 48 48 48 48 48 48 48
			2	20-perc	ent-m	olybder	num quate	rnary	alloys	₃ 8		
					Alpha	P	Mu	Alpha	P	Mu		
457899789901234545612 24559978990123555555555555555555555555555555555555	3 26 29 68 28 6 48 16 4 2 4 19 22 18 4 12 4	77 24 48 48 32 12.8 64 48 32 24 12.8 12.8 24 61.6 47.7	56.45 40.8 57.33 16.2 43.4 17.8 47.2 47.2 47.2 47.2 47.2 47.2 47.2 47.2	20.65 58 20.00 20.	100 90 100 100 95 100 	10	Trace	100 95 100 100 95 100 80 80 80 80 80 80 80 80 80 80 80 80 8	5	Trace	Alpha Alpha Alpha Alpha Alpha	泰安泰塔塔塔塔塔塔塔塔塔泰塔塔塔塔 泰

⁸All forged prior to annealing.

^bChemical composition from analysis. Composition given for other alloys is the intended melted analysis.

^cRemaining percentage is unidentified second phase.

dAlso trace of unidentified second phase.

TABLE VII—

TYPICAL X-RAY DIFFRACTION PATTERN OF FACE
CENTERED CUBIC ALPHA PHASE

Line	Estimated intensity	θ	đ	Radiation Cr K	hkl
1	Medium Strong Very weak Medium Very weak Weak Weak Very weak	29.86	2.089	Beta	111
2		33.17	2.089	Alpha	111
3		35.21	1.804	Beta	200
4		39.30	1.805	Alpha	200
5		54.84	1.272	Beta	220
6		63.85	1.273	Alpha _l	220
7		64.01	1.273	Alpha ₂	220
8		73.53	1.085	Beta	311



NACA TN 2683 43

TABLE VIII

TYPICAL X-RAY DIFFRACTION PATTERN OF BODY-

CENTERED CUBIC EPSILON PHASE

Line	Estimated intensity	θ	. ď	Radiation Cr K	hkl
12345678	Medium strong Strong Weak Weak Very weak Medium weak Strong Medium strong	30.69 34.13 46.43 52.75 52.87 62.64 77.42 77.89	2.039 2.037 1.435 1.435 1.435 1.171 1.171	Beta Alpha Beta Alpha ₁ Alpha ₂ Beta Alpha ₁ Alpha ₂	110 110 200 200 200 211 211 211



.

TABLE IX. - TYPICAL X-RAY DIFFRACTION PATTERNS OF

CHROMIUM-NICKEL-MOLYBDENUM AND CHROMIUM-

COBALT-MOLYBDENUM SIGMA PHASES

Line	Estimated intensity	в	đ	Radiation Cr K	Line	Estimated intensity	Ð	đ	Radiation Cr K
		Cr-N1-1	ngia oN	a phase, d	eterm.	ined from alloy	587		
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17	Medium Medium weak Medium weak Medium weak Medium weak Medium Medium weak Medium Medium weak Medium Medium Medium strong Strong Strong Medium Very weak Very very weak	26.66 27.38 28.25 28.61 29.51 30.40 31.25 31.76 32.87 33.75 34.81 35.60 36.58 37.78	2.172 2.160 1.917 2.057 2.002 1.964 1.918 1.864	Alpha Beta Alpha Alpha Alpha Beta Beta Alpha	20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36	Very weak Very very weak Weak Medium Medium weak Weak Very very weak Weak Very weak Weak Very weak Medium weak Weak Weak Very weak Weak Very weak Very weak	58.14 61.04 63.52 63.72 64.77 64.94 65.57 65.72 65.95 66.15 66.86 67.03 68.74 68.95 69.67	1.347 1.347 1.306 1.276 1.263 1.263 1.255 1.255 1.251 1.243 1.243 1.226 1.226 1.218	Alpha Alpha
18	Very very weak Very very weak Medium weak	39.70	1.789 1.672	Alpha Alpha	37 38	Very weak Very weak Very very weak	74.68	1.185	Alpha ₁ Alpha ₂
-	<u> </u>	Cr-Co-N	io sigm	a phase, d	eterm	ined from allöy	530		
1	Very weak	$\overline{}$	2.651	Alpha	21	Very weak		1.343	Alphal
2	Very weak		2.314	Beta	22	Very very weak		1.343	Alpha
3	Weak		2.473	Alpha	23	Medium strong		1.273	Alpha ₁
4	Medium weak		2.410	Alpha	24_	Medium		1.273	Alpha
5	Medium	28.68		Alpha	25	Medium		1.258	Alpha
6	Medium	29.58		Alpha	26	Medium weak	65.46		Alpha
7	Weak	30.44		Beta	27	Medium strong	65.89		Alpha
8	Medium weak	31.35		Beta	28	Medium weak	66.05		Alpha ₂
9	Medium strong	31.88		Alpha	29	Weak	66.42		Alpha ₁
10	Medium	32.11		Alpha	30	Very weak	66.65		Alpha
111	Medium strong	32.95	-	Beta	31	Medium weak	67.23		Alpha ₁
12	Medium	33.85		Alpha	32	Medium weak	67.42		Alpha
13	Strong	34.86		Alpha	33	Weak	67.71	- 1	Alpha ₁
14	Strong	35.69		Alpha	34	Very weak	67.95		Alpha
15	Strong	36.65		Alpha	_	Medium strong	69.31		Alpha ₁
16	Medium	37.87		Alpha	36	Medium	69.6	1.221	Alpha
17	Medium weak	39.81		Alpha	37	Weak	70.4	1.213	Alpha ₁
18	Very very weak			Alpha	38	Weak	70.63		Alpha
19	Very weak	55.16		Alpha	39	Weak	75.44	1	Alpha
20	Very weak	55.33		Alpha	49	Very weak	75.73		Alpha

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TABLE X

TYPICAL X-RAY DIFFRACTION PATTERN OF DELTA PHASE

DETERMINED FROM ALLOY 522

Line	Estimated intensity	θ	đ	Radiation Cr K	Line	Estimated intensity	θ	đ	Radiation Cr K
1	Weak	25.24	2.679	Alpha	23	Very weak	38.24	1.847	Alpha
2	Very weak	25.46	2.660	Alpha	24	Very very weak	38.91	1.820	Alpha
3 4	Weak	28.09	2.428	Alpha	25	Very very weak	40.85	1.748	Alpha
	Weak	28.27	2.196	Beta	26	Very weak	57.77	1.351	Alpha ₁
5	Very very weak	28.64	2.384	Alpha	27	Very very weak	57.89	1.351	Alpha ₂
6	Very weak	29.03	2.144	Beta	28	Very weak	58,90	1.334	Alphal
7	Weak	29.90	2.086	Beta	29	Very very weak	59.08	1.334	Alpha
8	Medium weak	30.37	2.058	Beta	30	Weak	63.94	1.272	Alpha
9	Weak	30.70	2.038	Beta	31	Very weak	64.15	1.272	Alpha
10	Strong	31.23	2.204	Alpha	32	Very weak	65.85	1.252	Alpha
11	Weak	31.42	2.192	Alpha	33	Very very weak	65.99	1.253	Alpha
12	Very weak	31.62	1.983	Beta	34	Very weak	66.36	1.247	Alpha ₁
13 14	Very very weak	32.01	2.156	Alpha	35	Very very weak	66.52	1.248	Alpha
14	Medium weak	32.25	2.142	Alpha	36	Weak	66.73	1.244	Alpha
15	Very very weak	32.77	2.113	Alpha	37	Very weak	66.93	1.244	Alpha
16	Very very weak	33.01	1.910	Beta	38	Very very weak	67.51	1.236	Alphal
17	Medium strong	33.24	2.084	Alpha	39	Very weak	67.74	1.237	Alpha
18	Strong	33.77	2.056	Alpha	40	Weak	68.02	1.232	Alphaj
19	Medium	34.14	2.037	Alpha	41	Very weak	68.22	1.232	Alpha
20	Medium	35.19	1.983	Alpha	142	Weak	70.64	1.211	Alpha ₁
21	Medium weak	36.35	1.929	Alpha	43	Weak	70.89	1.211	Alpha
22	Very weak	36.72	1.912	Alpha	44	Very weak	74.01	1.189	Alpha ₁
		•			45	Very very weak	74.38	1.188	Alphaz

TABLE XI

TYPICAL X-RAY DIFFRACTION PATTERN OF MU PHASE

DETERMINED FROM ALLOY 678

Line	Estimated intensity	θ	đ	Radiation Cr K	Line	Estimated intensity	θ	đ	Radiation Cr K
1 2 3 4	Medium weak Very weak Medium strong Very weak	26.00 28.65 28.83 29.26	2.608 2.169 2.370 2.339	Alpha Beta Alpha Alpha	17 18 19 20	Weak Very weak Very very weak Very weak	36.92 38.98 39.76 56.73	1.903 1.817 1.787 1.366	Alpha Alpha Alpha Alpha
5 6	Very very weak Medium weak	29.71 30.19	2.306 2.069	Alpha Beta	21 22	Very very weak Very weak	56.91 59.64	1.366	Alpha ₂ Alpha ₁
7	Very very weak	30.57	2.045	Beta	23	Very very weak	59.80	1.324	Alpha2
8	Weak	31.00	2.020	Beta	24	Weak	61.43	1.301	Alphal
9	Medium	31.83	2.166	Alpha	25	Very weak	61.62	1.301	Alpha ₂
10	Medium	32.52	2.125	Alpha	26	Weak	62.51	1.288	Alpha _l
11	Very very weak	33.17	1.901	Beta	27	Very weak	62.68	1.288	Alpha
12	Strong	33,52	2.070	Alpha	28,	Medium weak	70.04	1.216	Alpha ₁
13	Medium weak	33.98	2.045	Alpha	29	Weak	70.26	1.216	Alpha
14	Medium	34.48	2.019	Alpha	30	Medium weak	74.83	1.184	Alpha
15	Very very weak	34.91	1.998	Alpha	31	Weak	75.19	1.184	Alpha
16	Very weak	35.83	1.952	Alpha					



TABLE XII

TYPICAL X-RAY DIFFRACTION PATTERN OF P PHASE

DETERMINED FROM ALLOY 580

Line	Estimated intensity	θ	đ	Radiation Cr K	Line	Estimated intensity	θ	d.	Radiation Cr K
1	Very weak	25.54	2.650	Alpha	25	Medium weak	35.14	1.985	Alpha
2	Very very weak	25.82	2.388	Beta	26	Weak	36.44	1.925	Alpha
3	Very very weak	25.9	2.381	Beta	27	Weak	36.76	1.910	Alpha
4	Very very weak	26.51	2.329	Beta	28	Weak	37.07	1.897	Alpha
5	Weak	28.35	2.191	Beta	29	Very weak	38.69	1.829	Alpha
6	Medium weak	28.55	2.391	Alpha	30	Medium weak	60.2	1.317	Alpha
7	Medium weak	28.69	2.381	Alpha	31	Weak	60.39	1.317	Alpha
8	Medium weak	29.11	2,137	Beta	32	Weak	60.99	1.307	'Alpha _l
9	Weak	29.36	2.332	Alpha	33	Very weak	61.1	1.307	Alphae
10	Medium weak	29.65	2.102	Beta	34	Weak	61.5	1.300	Alpha
1,1	Weak	29.98	2.287	Alpha	35	Weak	61.62	1.301	Alpha
12	Weak	30.21	2.067	Beta	36	Weak	62.71	1.285	Alpha
13	Weak	30.65	2.040	Beta	37	Very weak	62.88	1.286	Alpha
14	Weak	30.81	2.031	Beta	38 39	Medium weak	63.48	1.277	Alpha
15	Strong	31.40	2.190	Alpha	39	Medium weak	65.06	1.260	Alphal
16	Weak	31.7	2.175	Alpha	40	Weak	65.24	1.269	Alpha
17	Medium weak	32.32	2.137	Alpha	41	Medium weak	67.34	1.238	Alphal
18	Medium	32.93	2.102	Alpha	42	Weak	67.51	1.239	Alpha
19	Medium	32.97	2.101	Alpha	43	Medium strong	73.17	1.194	Alpha
20	Medium	33.36	2.078	Alpha	1414	Medium weak	73.47	1.194	Alpha
21	Medium	33.58	2.067	Alpha	45	Weak	74.01	1.189	Alpha _l
22	Medium weak	33.85	2.052	Alpha	46	Very weak	74.28	1.189	Alpha
23	Medium	34.04	2.041	Alpha	47	Medium weak	76.42	1.175	Alphal
24	Medium strong	34.26	2.031	Alpha	48	Weak	76.83	1.175	Alpha ₂

TABLE XIII

TYPICAL X-RAY DIFFRACTION PATTERN OF R PHASE

DETERMINED FROM ALLOY 592

Line	Estimated intensity	θ	đ	Radiation Cr K	Line	Estimated intensity	0	đ	Radiation Cr K
1 2 3 4 5 6 7 8 9	Weak Very strong Very strong Very weak Very very weak Weak Medium weak Medium weak Medium weak Medium weak	25.45 25.94 26.18 26.43 26.87 27.52 28.59 28.81 29.29 29.59	2.659 2.612 2.592 2.336 2.529 2.473 2.173 2.159 2.337 2.107	Alpha Alpha Alpha Alpha Beta Alpha Beta Beta Alpha Beta Beta	22 23 24 25 26 27 28 29 30 31	Medium strong Medium Very weak Weak Very weak Medium weak Very weak Very weak Wery weak Wery weak	35.53 35.81 37.22 37.42 38.20 40.24 62.20 62.22 63.23 63.42	1.966 1.954 1.890 1.881 1.849 1.770 1.291 1.293 1.280	Alpha
11 12	Weak Medium weak	29.91 30.45	2.291 2.053	Alpha Beta	32 33	Medium weak Weak	64.27 64.49	1.268	Alpha ₁ Alpha ₂
13 14 15 16 17 18 19	Very weak Weak Weak Strong Strong Medium strong Medium strong	30.71 31.27 31.51 31.76 31.97 32.87 33.85	2.238 2.004 2.186 2.171 2.159 2.106 2.052	Alpha Beta Alpha Alpha Alpha Alpha Alpha	34 35 36 37 38 39 40	Very weak Very weak Medium weak Weak Medium weak Weak Medium weak	64.81 66.51 66.69 68.18 68.39 69.62 69.90	1.263 1.246 1.246 1.231 1.231 1.219 1.219	Alphal Alphal Alphal Alphal Alphal Alphal
20 21	Strong Medium strong	34.76 35.11	2.005 1.987	Alpha Alpha	41 42	Weak Weak	71.24	1.207	Alpha ₂

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TABLE XIV

DETERMINATION OF ALPHA THREE-PHASE-FIELD CORNER

BY X-RAY DIFFRACTION IN COBALT-NICKEL-

MOLYBDENUM SYSTEM

Data for fig. 26

Alloy	Weight percent cobalt	Extrapolated value of a _o (kX)
399 552 408 401 679 627 486 464 551 442	77 64 50 30 15 10 0 Three-phase alloy Three-phase alloy At boundary of three-phase field	3.5925 3.6008 3.6028 3.6133 3.6235 3.6320 3.6351 3.6287 3.6300 3.6307



TABLE XV

DETERMINATION OF SOLUBILITY LIMIT IN

COBALT-NICKEL-MOLYBDENUM MU PHASE

BY X-RAY DIFFRACTION

Data for fig. 27

Alloy	Weight percent nickel	θ	$\sin heta$	đ
421 678 550 536 537 523 442 464 551	0 12 18.77 24.12 26 29 At boundary of three-phase field Three-phase Three-phase	70.45 69.94 69.59 69.47 69.46 69.45	0.94235 .93953 .93771 .93724 .93649 .93642 .93637	1.2124 1.2160 1.2184 1.2190 1.2199 1.2200 1.2202



TABLE XVI

DETERMINATION OF ALPHA THREE-PHASE-FIELD CORNERS

BY X-RAY DIFFRACTION IN CHROMIUM-COBALT-

MOLYBDENUM SYSTEM

Data for fig. 28

Alloy	Weight percent chromium	Extrapolated value of a _o (kX)
399 487 413 665 645 412 606 644 471 398	0 4 10 14.7 17.42 19.36 21.98 24.9 28.83 31.7	3.5925 3.5870 3.5834 3.5832 3.5855 3.5890 3.5886 3.5810 3.5771



TABLE XVII

DETERMINATION OF ALPHA THREE-PHASE-FIELD CORNERS

BY_X-RAY DIFFRACTION IN CHROMIUM-NICKEL-

MOLYBDENUM SYSTEM

Data for fig. 29

Alloy	Weight percent chromium	Extrapolated value of a _o (kX)
486	0	3.6351
688	5.5	3.6276
636	11.3	3.6245
460	14.3	3.6213
469	22.1	3.6189
457	23	3.6186
438	25.5	3.6162
435	36.25	3.6073
389	42.75	3.5956
614	43.5	3.5956

 8 From reference 1; 8 O redetermined in the present investigation.

TABLE XVIII

VALUES OF d OF CHROMIUM-NICKEL-MOLYBDENUM TERNARY

SIGMA PHASE AS FUNCTION OF CHROMIUM CONTENT

Data for fig. 30

Alloy	Weight percent chromium	d value ^l
509 508 641 482 603 483 604 447 491 586	25.06 32.0 22.76 28.0 36.0 33.0 45.0 40.0 45.0	1.2996 1.2969 1.2970 1.2928 1.2916 1.2857 1.2853 1.2832 1.2769 1.2768

¹d values determined from twenty-third line of sigma patterns.

TABLE XIX.- DATA USED FOR PLOTTING ACTUAL SOLUBILITY OF CHROMIUM IN ALPHA PHASE AT VARIOUS CONSTANT MOLYBDENUM CONTENTS AS FUNCTION OF ACTUAL COBALT CONTENT

Data for fig. 32

Weight percent molybdenum	Weight percent cobalt	Maximum weight percent chromium in alpha solid solution
0 0 0 0 0 0	0 10 18 20 30 40 50 60	42.2 44.3 44.5 43.4 40.8 38.6 37.0 35.0 34.0
2.5 5.5 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2	C 9.75 19.5 24.3 29.25 39.0 48.75 58.6 63.4	43.3 41.4 40.9 40.5 39.9 38.0 36.0 34.0 33.2
555555	0 9.5 19.0 28.5 43.5 57.0 64.5	42.2 40.0 39.4 37.6 35.0 32.0 30.5
10 10 10 10 10 10 10	0 9.0 18.0 34.2 45.0 48.6 54.0 64.4	36.9 33.75 31.5 29.25 27.9 27.0 26.6 25.7
20 20 20 20 20 20 20 20 20 20 20 20	0 12.8 24.0 32.0 48.0 56.0 60.0 61.5 64.0 72.0 76.0	24.0 19.2 17.6 15.6 12.0 11.2 10.8 10.5 9.2 5.6 4.0

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TABLE XX

DETERMINATION OF THREE-PHASE-FIELD CORNERS

ON 20-PERCENT-MOLYBDENUM QUATERNARY

ALPHA PHASE BOUNDARY

Data for fig. 33

Alloy	Actual weight percent cobalt	Extrapolated value of a _o (kX)
457	0	3.6186
524	12.2	3.6128
525	24.0	3.6097
526	32.0	3.6042
652	47.6	3.6004
651	61.6	3.5969
501	64.0	3.5900
487	76.0	3.5870



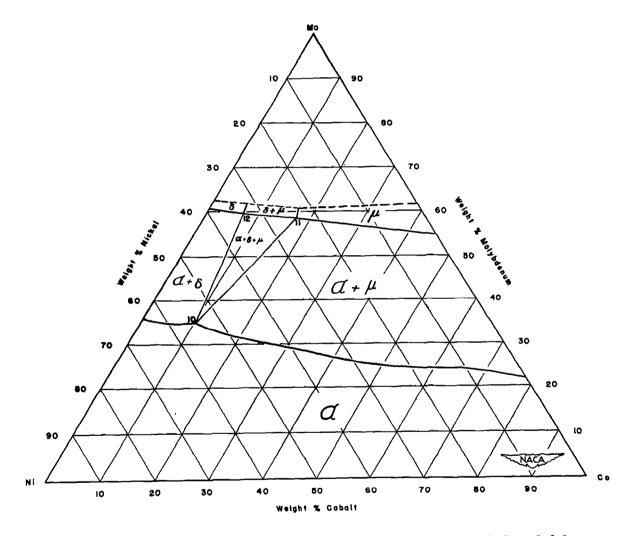


Figure 1.- The $1200^{\rm O}$ C isothermal section of cobalt-nickel-molybdenum ternary system.

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NACA TN 2683 57

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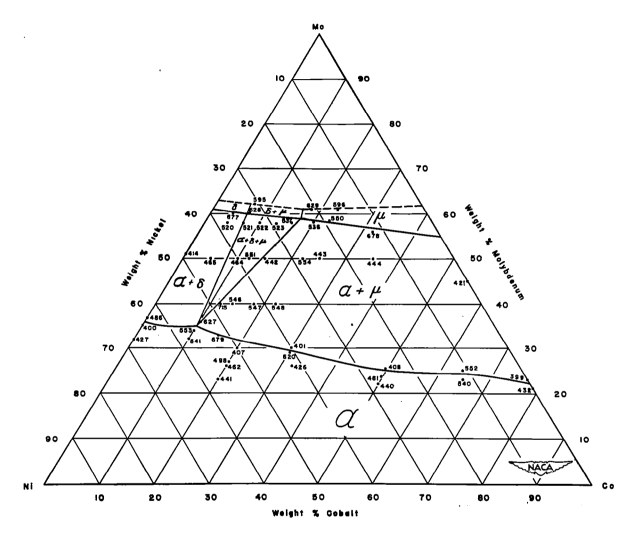


Figure 2.- The 1200° C isothermal section of cobalt-nickel-molybdenum ternary system with alloy compositions indicated.

58 NACA IN 2683

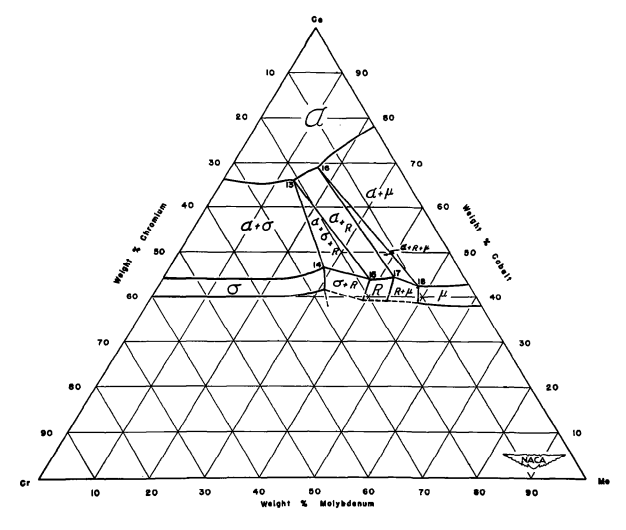


Figure 3.- The 1200° C isothermal section of chromium-cobalt-molybdenum ternary system.

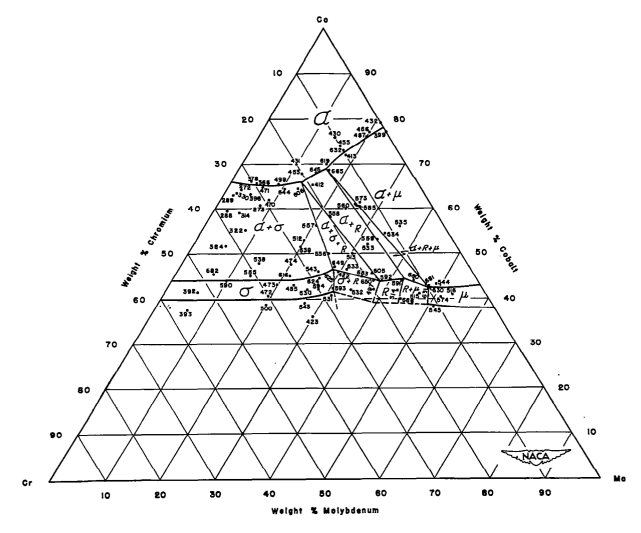


Figure 4.- The 1200°C isothermal section of chromium-cobalt-molybdenum ternary system with alloy compositions indicated.

60 NACA TN 2683

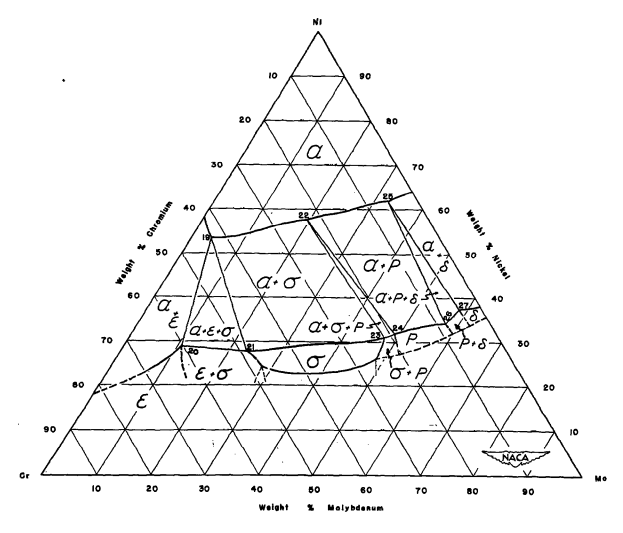


Figure 5.- The 1200° C isothermal section of chromium-nickel-molybdenum ternary system.

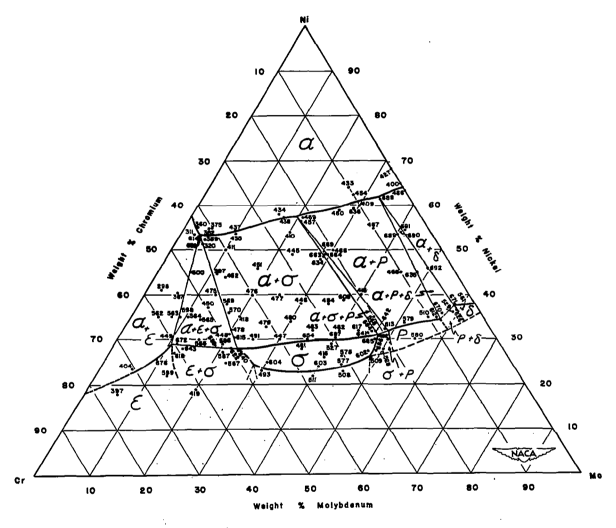


Figure 6.- The 1200°C isothermal section of chromium-nickel-molybdenum termary system with alloy compositions indicated.

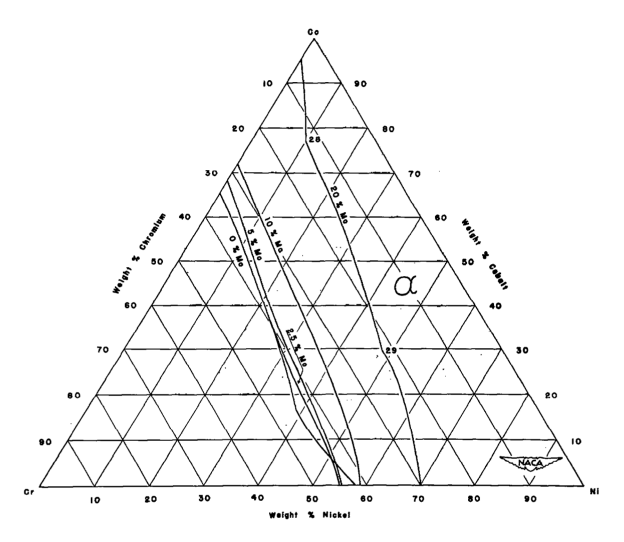


Figure 7.- Alpha-phase-field boundary at 1200°C in chromium-cobalt-nickel-molybdenum quaternary system for constant molybdenum contents of 2.5, 5, 10, and 20 percent.

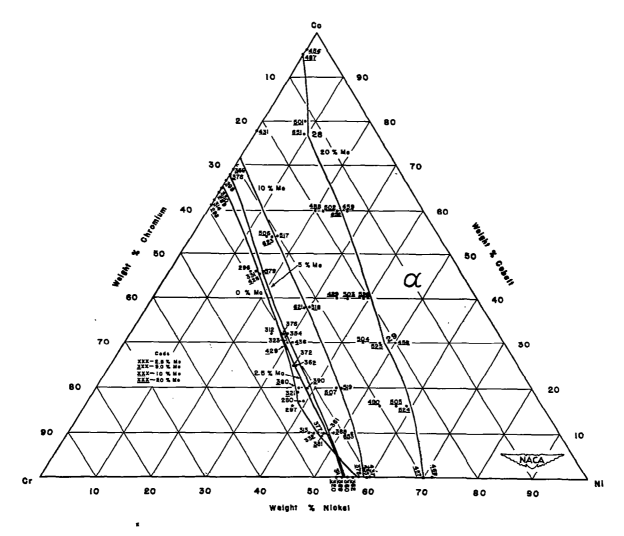


Figure 8.- Alpha-phase-field boundary at 1200°C in chromium-cobaltnickel-molybdenum quaternary system for constant molybdenum contents of 2.5, 5, 10, and 20 percent with alloy compositions indicated.

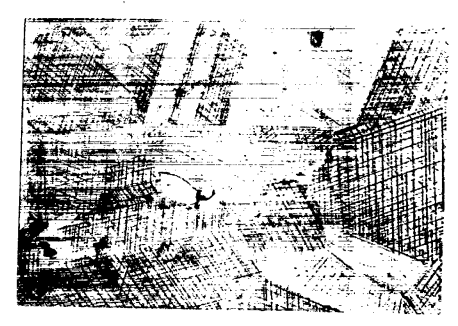


Figure 9.- Alloy 296 containing 38.02 percent chromium, 44.85 percent cobalt, 14.63 percent nickel, and 2.5 percent molybdenum. Etched according to procedure 1; small particles of second phase in an alpha matrix; transformation striations revealed in alpha phase; X500.

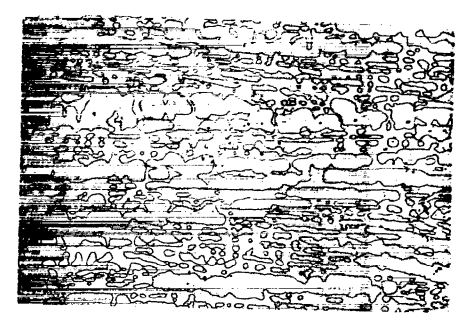


Figure 10.- Alloy 443 containing 25 percent cobalt, 25 percent nickel, and 50 percent molybdenum. Etched according to procedure 4 but stain omitted; alpha phase arranged in banded dendritic pattern in matrix of mu phase; X250.

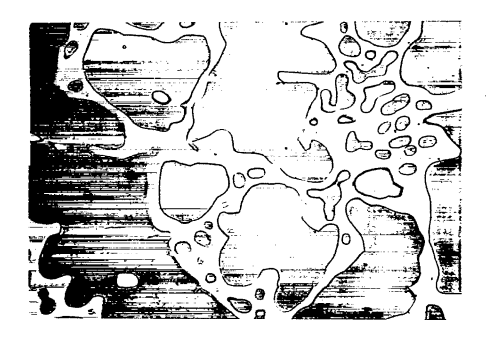


Figure 11.- Alloy 608 containing 23.5 percent chromium, 39 percent nickel, and 37.5 percent molybdenum. Etched and stained according to procedure 2; minor amounts of unstained alpha in matrix of stained sigma phase; X250.



Figure 12.- Alloy 467 containing 10 percent chromium, 55 percent nickel, and 35 percent molybdenum. Etched according to procedure 2 but stain omitted; typical distribution of minor amounts of P phase in matrix of alpha; X250.

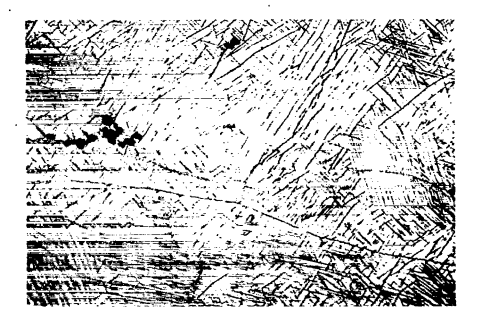


Figure 13.- Alloy 599 containing 65 percent chromium, 22 percent nickel, and 13 percent molybdenum. Etched and stained according to procedure 2; heavy Widmanstatten precipitate of sigma in epsilon phase; dark oxide inclusions and grain boundaries shown in epsilon phase; X500.

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Figure 14.- Alloy 600 containing 48.5 percent chromium, 45.4 percent nickel, and 6.1 percent molybdenum. Etched and stained according to procedure 2; heavily stained and severely cracked particles of sigma plus lightly stained epsilon with few cracks in matrix of unstained alpha phase; X250.

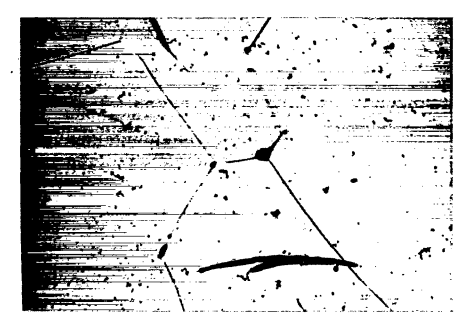


Figure 15.- Alloy 602 containing 25 percent chromium, 27.5 percent nickel, and 47.5 percent molybdenum. Etched and stained according to procedure 2; grain boundaries revealed in cracked matrix of sigma phase; X500.



Figure 16.- Alloy 492 containing 20 percent chromium, 30 percent nickel, and 50 percent molybdenum. Etched and stained according to procedure 2; particles of sigma phase clearly delineated in matrix of P phase; grain boundaries revealed; X1000.

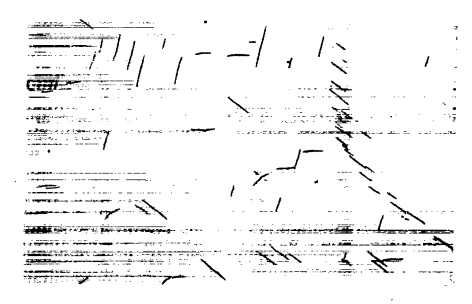


Figure 17.- Alloy 392 containing 52.6 percent chromium, 41.4 percent cobalt, and 6 percent molybdenum. Etched and stained according to procedure 3; scattered Widmanstatten precipitate in matrix of sigma phase; identity of precipitate unknown; X500.

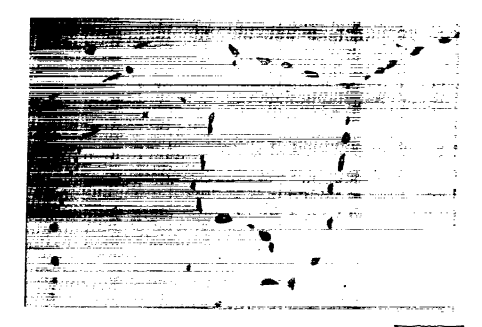


Figure 18.- Alloy 471 containing 28.83 percent chromium, 64.8 percent cobalt, and 6.37 percent molybdenum. Etched and stained according to procedure 3; small particles of stained sigma phase outlining grains of unstained alpha phase; X500.

NACA IN 2683 69

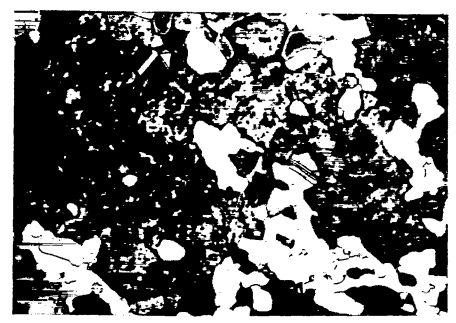


Figure 19.- Alloy 522 containing 10 percent cobalt, 32 percent nickel, and 58 percent molybdenum. Etched and stained according to procedure 4; minor amounts of unstained (white) alpha phase plus small grains of mu phase stained to various colors and showing occasional annealing twins in matrix of delta phase; delta phase unevenly attacked and heavily stained; X500.



Figure 20.- Alloy 626 containing 4 percent chromium, 36.5 percent nickel, and 59.5 percent molybdenum. Etched and stained according to procedure 2; traces of P phase in matrix of delta; grain boundaries in delta phase very faintly visible; X1500.

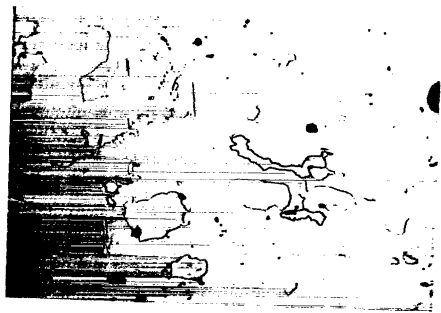


Figure 21.- Alloy 550 containing 22.74 percent cobalt, 18.77 percent nickel, and 58.49 percent molybdenum. Etched according to procedure 4 but stain omitted; structure of mu phase; preferential attack probably due to orientation effect: X500.



Figure 22.- Alloy 669 containing 22 percent chromium, 49 percent nickel, and 29 percent molybdenum. Etched and stained according to procedure 2; of alpha phase; slight contrast between sigma and P particles; X500.

71

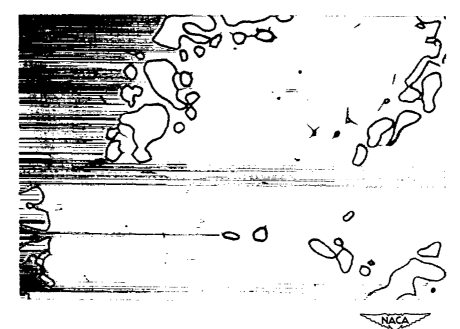


Figure 23.- Alloy 605 containing 18.5 percent chromium, 45.5 percent cobalt, and 36 percent molybdenum. Etched and stained according to procedure 3; small particles of stained sigma plus minor amounts of unstained (white) alpha phase in stained matrix of R phase; X500.

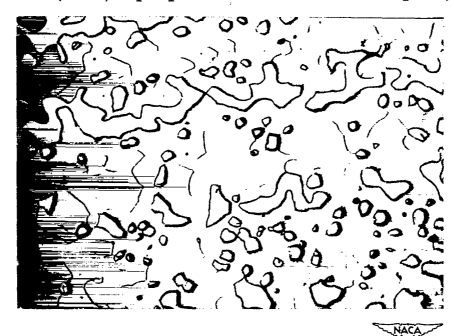


Figure 24.- Alloy 513 containing 19.88 percent chromium, 49.78 percent cobalt, and 30.34 percent molybdenum. Etched and stained according to procedure 3; phase contrast shown in three-phase alpha-sigma-Ralloy; alpha is unstained (white), sigma is stained (darkest), and R is lightly stained; X500.

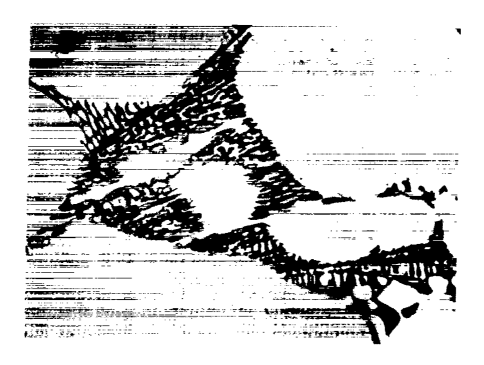


Figure 25.- Alloy 376 containing 38.76 percent chromium, 31.2 percent cobalt, 27.54 percent nickel, and 2.5 percent molybdenum. Etched lightly according to procedure 1; dark etching zirconium impurity phase associated with small second-phase particles of sigma in matrix of alpha phase; X2000.

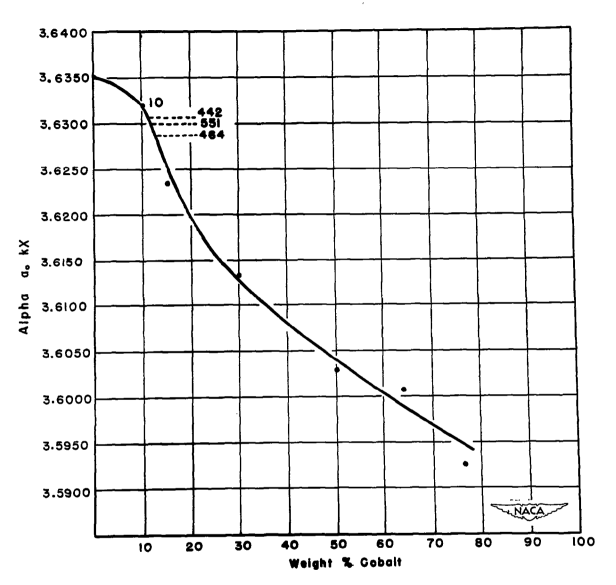


Figure 26.- Variation of lattice parameter at 1200°C of alpha boundary alloys in cobalt-nickel-molybdenum ternary system plotted as function of cobalt content.

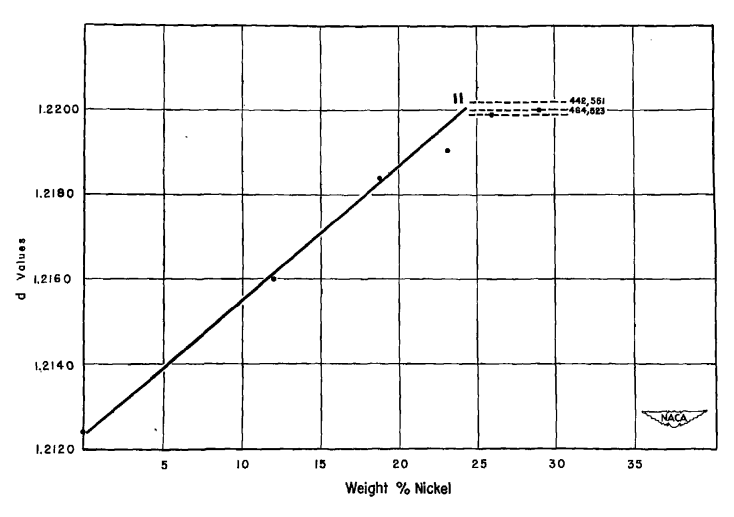


Figure 27.- Variation at 1200° C of d value of twenty-eight X-ray diffraction line of mu phase in cobalt-nickel-molybdenum ternary system plotted as function of nickel content.

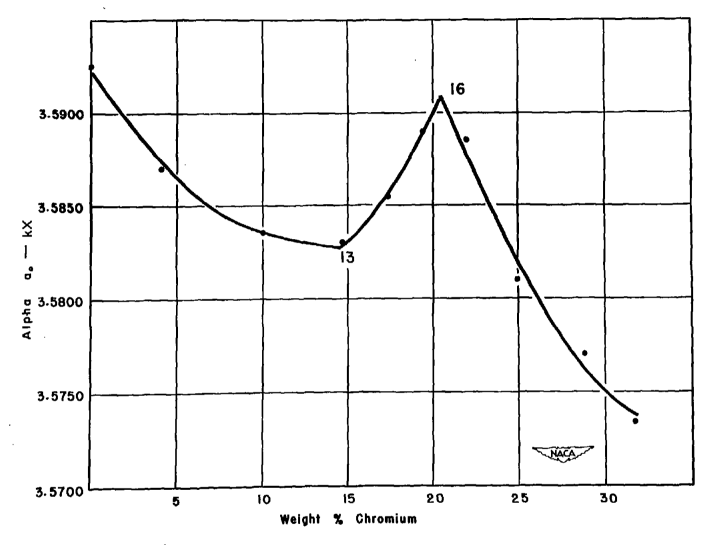


Figure 28. - Variation of lattice parameter at 1200°C of alpha boundary alloys in chromium-cobalt-molybdenum ternary system plotted as function of chromium content.

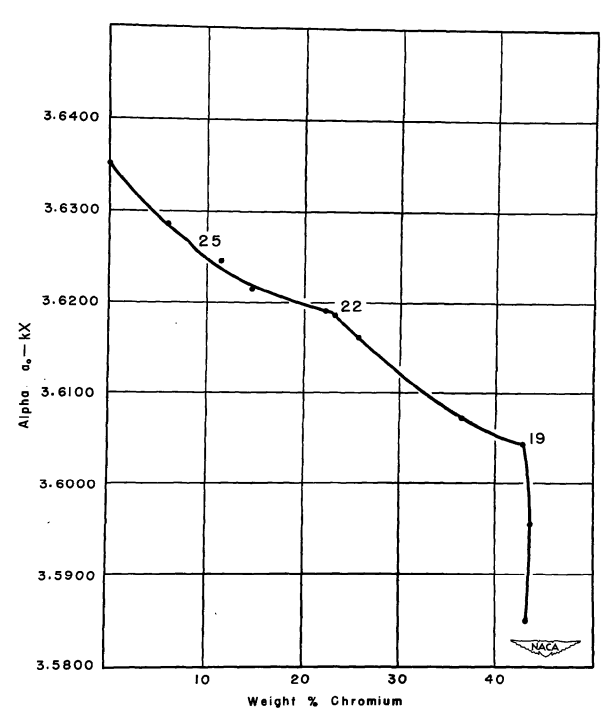


Figure 29.- Variation of lattice parameter at 1200° C of alpha boundary alloys in chromium-nickel-molybdenum ternary system plotted as function of chromium content.

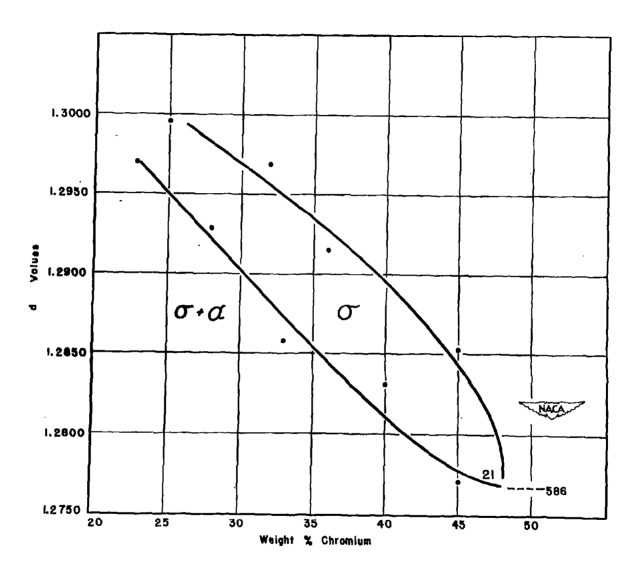


Figure 30.- Variation at 1200°C of d values of twenty-third X-ray diffraction line of sigma phase in chromium-nickel-molybdenum ternary system plotted as function of chromium content.

78

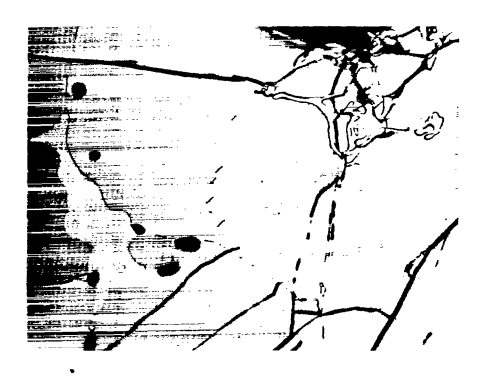


Figure 31.- Alloy 447 containing 40 percent chromium, 30 percent nickel, and 30 percent molybdenum. Etched and stained according to procedure 2; small particles of phase not identifiable from figure 6 plus unstained alpha in matrix of sigma phase; X750.

NACA TN 2683

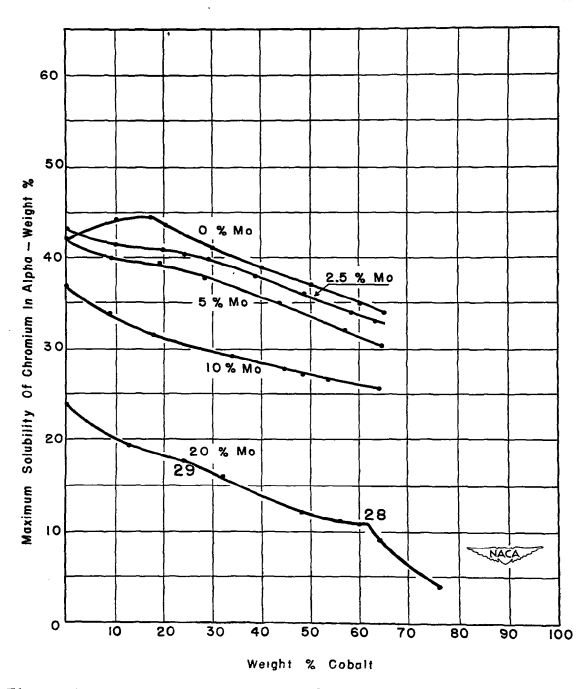


Figure 32.- Maximum solubility at 1200°C of chromium in alpha phase of chromium-cobalt-nickel-molybdenum alloys plotted as functions of actual cobalt content for constant molybdenum contents of 2.5, 5, 10, and 20 percent.

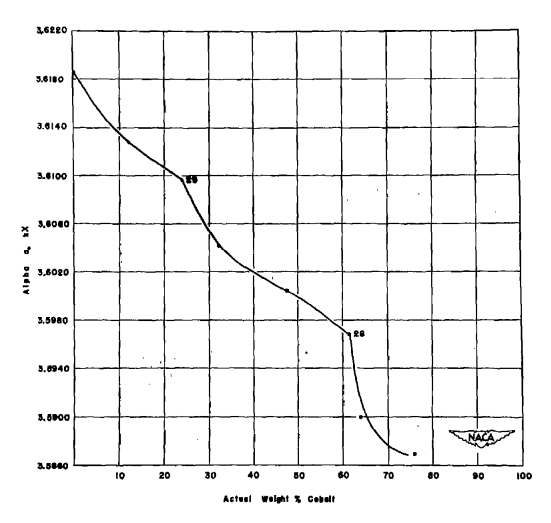


Figure 33.- Variation of lattice parameter at 1200°C of alpha boundary alloys in 20-percent-molybdenum section of chromium-cobalt-nickelmolybdenum quaternary system plotted as function of actual cobalt content.

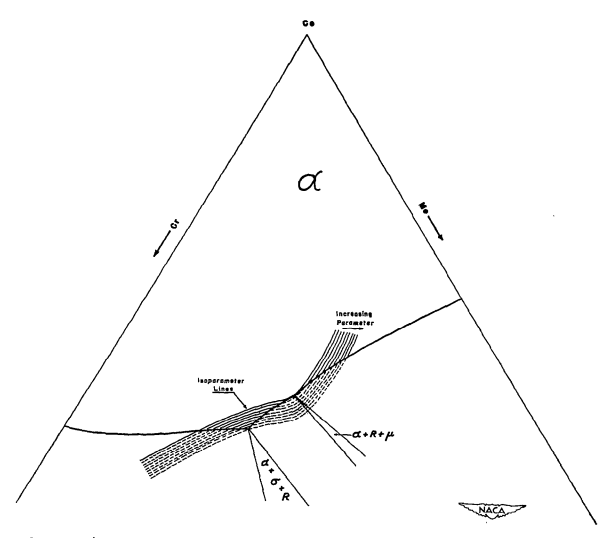


Figure 34.- Schematic sketch of alpha-phase-field boundary at 1200°C in chromium-cobalt-molybdenum ternary system with hypothetical isoparameter lines shown.